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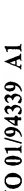
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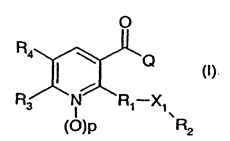
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(54) Title: SUBSTITUTED PYRIDINE HERBICIDES





(57) Abstract: Compounds of the formula (I) in which the substituents are as defined in claim 1 are suitable for use as herbicides.

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SUBSTITUTED PYRIDINE HERBICIDES

The present invention relates to novel herbicidally active pyridine ketones, to processes for their preparation, to compositions which comprise these compounds, and to their use for controlling weeds, in particular in crops of useful plants, or for inhibiting plant growth.

Pyridine ketones having herbicidal action are described, for example, in WO 00/15615 and WO/0039094.

We have now found novel pyridine ketones having herbicidal and growth-inhibiting properties.

The present invention thus provides compounds of the formula I

in which

p is 0 or 1;

 R_1 is a C_1 - C_6 alkylene, C_3 - C_6 alkenylene or C_3 - C_6 alkynylene chain which may be mono- or polysubstituted by halogen or R_5 , where the unsaturated bonds of the chain are not attached directly to the substituent X_1 ;

 X_1 is oxygen, -O(CO)-, -(CO)O-, -O(CO)O-, -N(R₆)-O-, -O-NR₅₁-, thio, sulfinyl, sulfonyl, -SO₂NR₇-, -NR₅₂SO₂- or -NR₈-;

 R_2 is a C_1 - C_6 alkyl, C_3 - C_6 alkenyl or C_3 - C_6 alkynyl group which is mono- or polysubstituted by halogen, hydroxyl, amino, formyl, nitro, cyano, mercapto, carbamoyl, C_1 - C_6 alkoxy, C_1 - C_6 alkoxycarbonyl, C_2 - C_6 alkenyl, C_2 - C_6 haloalkenyl, C_2 - C_6 alkynyl, C_2 - C_6 haloalkynyl, C_3 - C_6 cycloalkyl, by halogen-substituted C_3 - C_6 cycloalkyl, or by C_3 - C_6 alkenyloxy, C_3 - C_6 alkynyloxy, C_1 - C_6 alkoxy, C_3 - C_6 haloalkenyloxy, cyano- C_1 - C_6 alkoxy, C_1 - C_6 alkoxy- C_1 - C_6 alkoxy- C_1 - C_6 alkoxy- C_1 - C_6 alkoxy, C_1 - C_6 alkoxy,

 C_1 - C_6 alkylsulfonyl, C_1 - C_6 alkylcarbonyl, C_1 - C_6 alkylsulfinyl, C_1 - C_6 alkylsulfonyl, C_1 - C_6 haloalkylthio, C_1 - C_6 haloalkylsulfinyl, C_1 - C_6 haloalkylsulfonyl, oxiranyl, which for its part may be substituted by C_1 - C_6 alkyl, or by (3-oxetanyl)oxy, which for its part may be substituted by C_1 - C_6 alkyl, or by benzylthio, benzylsulfinyl, benzylsulfonyl, C_1 - C_6 alkylamino, $di(C_1$ - C_6 alkyl)amino, $R_9S(O)_2O$, $R_{10}N(R_{11})SO_2$ -, thiocyanato, phenyl, phenoxy, phenylthio, phenylsulfinyl or phenylsulfonyl;

where the phenyl- or benzyl-containing groups for their part may be substituted by one or more C_1 - C_6 alkyl, C_1 - C_6 haloalkyl, C_1 - C_6 alkoxy, C_1 - C_6 haloalkoxy, halogen, cyano, hydroxyl or nitro groups, or

 R_2 is phenyl which may be mono- or polysubstituted by C_1 - C_6 alkyl, C_1 - C_6 haloalkyl, C_1 - C_6 alkoxy, C_1 - C_6 haloalkoxy, halogen, cyano, hydroxyl or nitro; or

 R_2 is C_3 - C_6 cycloalkyl, C_1 - C_6 alkoxy- or C_1 - C_6 alkyl-substituted C_3 - C_6 cycloalkyl, 3-oxetanyl or C_1 - C_6 alkyl-substituted 3-oxetanyl;

or, if Q is Q_2 or Q_3 , or is Q_1 in which R_{14} and R_{22} are a C_2 - C_3 alkylene chain, R_2 is additionally also a five- to ten-membered monocyclic or fused bicyclic ring system which may be aromatic, saturated or partially saturated and may contain 1 to 4 heteroatoms selected from the group consisting of nitrogen, oxygen and sulfur, where the ring system is attached directly or via a C_1 - C_4 alkylene, C_2 - C_4 alkylene, C_2 - C_4 alkylene, C_2 - C_4 alkylene, -N(R₁₂)-C₁-C₄alkylene, -SO-C₁-C₄alkylene or -SO₂-C₁-C₄alkylene group to the substituent X_1 and where each ring system may not contain more than two oxygen atoms and not more than two sulfur atoms and the ring system for its part may be mono-, di- or trisubstituted by C_1 - C_6 alkyl, C_1 - C_6 haloalkyl, C_2 - C_6 alkenyl, C_2 - C_6 haloalkynyl, C_2 - C_6 haloalkynyl, C_1 - C_6 alkoxy, hydroxyl, C_1 - C_6 haloalkoxy, C_3 - C_6 alkenyloxy, C_3 - C_6 alkynyloxy, mercapto, C_1 - C_6 alkylthio, C_1 - C_6 haloalkylthio, C_3 - C_6 alkenylthio, C_3 - C_6 haloalkenylthio, C_3 - C_6 alkynylthio, C_2 - C_5 alkoxyalkylthio, C_3 - C_5 acetylalkylthio, C_3 - C_6 alkoxycarbonylalkylthio, C_2 - C_4 cyanoalkylthio, C_1 - C_6 alkylsulfinyl, C_1 - C_6 haloalkylsulfinyl, C_1 - C_6 alkylsulfonyl, C_1 - C_6 haloalkylsulfonyl, aminosulfonyl, C₁-C₂alkylaminosulfonyl, di(C₁-C₂alkyl)aminosulfonyl, di(C₁-C₄alkyl)amino, halogen, cyano, nitro, phenyl and benzylthio, where phenyl and benzylthio for their part may be substituted on the phenyl ring by C_1 - C_3 alkyl, C_1 - C_3 haloalkyl, C_1 - C_3 alkoxy, C₁-C₃haloalkoxy, halogen, cyano or nitro, and where the substituents on the nitrogen in the

heterocyclic ring are different from halogen; or R₂ is hydrogen or unsubstituted C₁-C₈alkyl if

- a) R₁ is substituted by the group R₅, or
- b) Q is the group Q2, or

c) Q is the group Q_3 in which X_1 is -O(CO)-, -(CO)O-, -N(R₆)-O-, -O-NR₅₁-, -SO₂NR₇-, -NR₅₂SO₂- or -NR₈-; or

d) Q is the group Q_1 in which X_1 is $-N(R_6)-O_7$, $-O-NR_{51}$, $-SO_2NR_{77}$, $-NR_{52}SO_2$ - or $-NR_{87}$, or

e) Q is the group Q_1 in which R_{14} and R_{22} in Q_1 are a C_2 - C_3 alkylene chain and X_1 is -O(CO)-or -(CO)O-,

R₃ is C₁-C₃haloalkyl;

 R_4 is hydrogen, halogen, C_1 - C_3 alkyl, C_1 - C_3 alkoxy, C_1 - C_3 alkoxy- C_1 - C_3 alkoxy; C_1 - C_3 alkoxy;

 R_5 is hydroxyl, C_1 - C_6 alkoxy, C_3 - C_6 cycloalkyloxy, C_1 - C_6 alkoxy- C_1 - C_1 -

 R_6 , R_7 , R_8 , R_9 , R_{10} , R_{11} , R_{12} , R_{51} and R_{52} independently of one another are hydrogen, C_1 - C_6 alkyl, C_1 - C_6 alkyl, C_1 - C_6 alkoxy- C_1 - C_6 alkoxy, benzyl or phenyl, where phenyl and benzyl for their part may be mono- or polysubstituted by C_1 - C_6 alkyl, C_1 - C_6 haloalkyl, C_1 - C_6 alkoxy, C_1 - C_6 haloalkoxy, halogen, cyano, hydroxyl or nitro; where R_6 and R_9 are not simultaneously hydrogen and hydrogen, C_1 - C_6 alkoxycarbonyl or C_1 - C_6 alkylcarbonyl, respectively;

Q is Q₁

in which

 A_1 is $C(R_{14}R_{15})$, NR_{16} or oxygen;

 A_2 is $C(R_{17}R_{18})$, C(O), $-C=N-O-R_{19}$, oxygen, thio, sulfinyl, sulfonyl, $-NR_{20}$ or ethylene; with the provisos that A_1 is different from oxygen if A_2 is oxygen, C(O), thio, sulfinyl, $-C=N-O-R_{19}$, NR_{20} or $C(R_{17}R_{18})$, where R_{17} and R_{18} independently of one another are C_1-C_4 alkoxy, C_1-C_4 alkylsulfinyl, C_1-C_4 alkylsulfonyl; and A_1 is different from NR_{16} if A_2 is thio, sulfinyl or $C(R_{17}R_{18})$, where R_{17} and R_{18} independently of one another are C_1-C_4 alkoxy, C_1-C_4 alkylsulfinyl, C_1-C_4 alkylsulfonyl; R_{14} and R_{22} independently of one another are hydrogen, C_1-C_4 alkyl, C_1-C_4 alkylsulfonyl, C_3-C_4 alkenyl, C_3-C_4 alkynyl, C_1-C_4 alkylthio,

C₁-C₄alkylsulfinyl, C₁-C₄alkylsulfonyl, C₁-C₄alkylsulfonyloxy, C₁-C₄alkoxy,

C₁-C₄alkoxycarbonyl or C₁-C₄alkylcarbonyl;

 R_{15} and R_{21} independently of one another are hydrogen, C_1 - C_4 alkyl, C_1 - C_4 haloalkyl, C_3 - C_4 alkenyl or C_3 - C_4 alkynyl;

 R_{17} is hydrogen, $C_1\text{-}C_4$ alkyl, $C_1\text{-}C_4$ haloalkyl, $C_1\text{-}C_4$ alkoxy, $C_1\text{-}C_4$ alkylsulfinyl or $C_1\text{-}C_4$ alkylsulfonyl;

 R_{18} is hydrogen, C_1 - C_4 alkyl, C_1 - C_4 haloalkyl, C_3 - C_4 alkenyl, C_3 - C_4 alkynyl, C_1 - C_4 alkylsulfinyl, C_1 - C_4 alkylsulfonyl or C_1 - C_4 dialkoxyalkyl- C_1 - C_4 alkyl; R_{20} is C_1 - C_4 alkyl, C_3 - C_6 cycloalkyl, C_3 - C_4 alkenyl, C_3 - C_4 alkynyl, C_1 - C_4 alkylcarbonyl, C_1 - C_4 alkylcarbonyl or benzyl, where the phenyl group may be monoor polysubstituted by C_1 - C_6 alkyl, C_1 - C_6 haloalkyl, C_1 - C_6 alkoxy, C_1 - C_6 haloalkoxy, halogen, cyano, hydroxyl or nitro;

 R_{19} and R_{16} independently of one another are hydrogen, C_1 - C_4 alkyl, C_3 - C_6 cycloalkyl, C_3 - C_4 alkenyl, C_3 - C_4 alkynyl, benzyl or phenyl, where phenyl and benzyl for their part may be mono- or polysubstituted by C_1 - C_6 alkyl, C_1 - C_6 haloalkyl, C_1 - C_6 alkoxy, C_1 - C_6 haloalkoxy, halogen, cyano, hydroxyl or nitro;

or R₁₄ and R₂₂ together form a C₂-C₃alkylene chain;

or R_{14} and R_{15} together and/or R_{17} and R_{18} together and/or R_{21} and R_{22} together form a C_2 - C_4 alkylene chain which may be interrupted by oxygen and/or carbonyl and/or sulfur, with the proviso that the oxygen and sulfur atoms are separated by at least one methylene group; or R_{14} and R_{18} together form a C_2 - C_4 alkylene chain; or

R₂₂ and R₁₈ together form a C₂-C₄alkylene chain;

or R₁₈ together with R₂₂ or R₁₄ forms a direct bond;

or R₁₆ and R₁₈ together form a C₂-C₄alkylene chain;

 R_{13} is hydroxyl, O⁻M⁺, where M⁺ is an alkali metal cation or ammonium cation, halogen, $C_1\text{-}C_{12}$ alkylsulfonyloxy, amino, $C_1\text{-}C_4$ alkylthio, $C_1\text{-}C_{12}$ alkylsulfinyl, $C_1\text{-}C_{12}$ alkylsulfonyl, $C_1\text{-}C_{12}$ haloalkylsulfinyl, $C_1\text{-}C_{12}$ haloalkylsulfonyl, $C_1\text{-}C_6$ alkoxy- $C_1\text{-}C_6$ alkoxy- $C_1\text{-}C_6$ alkylsulfinyl, $C_1\text{-}C_6$ alkoxy- $C_1\text{-}C_6$ alkylsulfonyl, $C_3\text{-}C_{12}$ alkenylsulfinyl, $C_3\text{-}C_{12}$ alkenylsulfonyl, $C_3\text{-}C_1$ alkynylsulfinyl, $C_3\text{-}C_1$ alkynylsulfinyl, $C_3\text{-}C_1$ alkoxycarbonyl- $C_1\text{-}C_4$ alkoxycarbonyl- $C_1\text{-}C_4$ alkoxycarbonyl- $C_1\text{-}C_4$ alkoxycarbonyl- $C_1\text{-}C_4$ alkoxylsulfonyl, $C_1\text{-}C_4$ alkoxylsulfonyloxylsu

carbonyloxy, C_1 - C_{12} alkylthiocarbamoyl, where the alkyl, alkenyl and alkynyl groups may be substituted by halogen, C_1 - C_6 alkoxy, C_1 - C_6 alkylthio, C_1 - C_6 alkylsulfinyl, C_1 - C_6 alkylsulfonyl or cyano;

or R₁₃ is phenoxy, phenylthio, phenylsulfinyl, phenylsulfonyl, phenylsulfonylamino, phenylsulfonyloxy or benzoyloxy, where the phenyl groups for their part may be substituted by one or more halogen, nitro, cyano, C₁-C₄alkyl, C₁-C₄haloalkyl, C₁-C₄alkoxy or C₁-C₄haloalkoxy groups;

or R₁₃ is a group Het₁-thio, Het₂-sulfinyl, Het₃-sulfonyl, Het₄-(CO)O or Het₅-N(R₃₃); in which Het₁, Het₂, Het₃, Het₄ and Het₅ independently of one another are a five- to ten-membered monocyclic or fused bicyclic ring system which may be aromatic or partially saturated and may contain 1 to 4 heteroatoms selected from the group consisting of nitrogen, oxygen and sulfur, and where each ring system may not contain more than 2 oxygen atoms and not more than 2 sulfur atoms, and where the ring system itself can be substituted by C₁-C₆alkyl, C₁-C₆haloalkyl, C₁-C₆alkoxy, C₁-C₆haloalkoxy, C₁-C₆alkylthio, C₁-C₆alkylsulfinyl, C₁-C₆alkylsulfonyl, di(C₁-C₄alkyl)aminosulfonyl, di(C₁-C₄alkyl)amino, halogen, cyano, nitro or phenyl, and where the substituents on the nitrogen in the heterocyclic ring are different from halogen;

 R_{23} , R_{24} , R_{25} , R_{26} , R_{27} , R_{28} , R_{29} , R_{30} and R_{33} independently of one another are hydrogen or C_1 - C_6 alkyl;

or R_{23} and R_{24} together or R_{25} and R_{26} together or R_{27} and R_{28} together or R_{29} and R_{30} together are pyrolidino, piperidino, morpholino, thiomorpholino, which may be monopolysubstituted by methyl groups;

or Q is Q2

in which

 R_{34} is hydrogen, C_1 - C_4 alkyl, C_1 - C_4 haloalkyl, C_3 - C_6 cycloalkyl, C_2 - C_4 alkenyl, C_2 - C_4 alkynyl or benzyl, where the phenyl group may be mono- or polysubstituted by C_1 - C_6 alkyl, C_1 - C_6 haloalkoxy, halogen, cyano, hydroxyl or nitro;

 R_{35} is hydrogen, C_1 - C_4 alkyl, C_1 - C_4 haloalkyl, C_3 - C_6 cycloalkyl, C_3 - C_4 alkenyl, C_3 - C_4 alkynyl or benzyl, where the phenyl group may be mono- or polysubstituted by C1-C6alkyl, C1-C6haloalkyl, C₁-C₆alkoxy, C₁-C₆haloalkoxy, halogen, cyano, hydroxyl or nitro; R₃₆ is hydroxyl, O⁻M⁺, where M⁺ is an alkali metal cation or ammonium cation, halogen, C₁-C₁₂alkylsulfonyloxy, amino, C₁-C₄alkylthio, C₁-C₁₂alkylsulfinyl, C₁-C₁₂alkylsulfonyl, C₁-C₁₂haloalkylthio, C₁-C₁₂haloalkylsulfinyl, C₁-C₁₂haloalkylsulfonyl, C₁-C₆alkoxy- $C_1-C_6 alkylsulfinyl,\ C_1-C_6 alkylsulfinyl,\ C_1-C_6 alkylsulfonyl,\ C_3-C_{12} alkenyl-C_6 alkylsulfonyl,\ C_3-C_{12} alkenyl-C_6 alkylsulfonyl,\ C_8-C_{12} alkenyl-C_8 alkylsulfonyl,\ C_8-C_8 al$ thio, C_3 - C_{12} alkenylsulfinyl, C_3 - C_{12} alkenylsulfonyl, C_3 - C_{12} alkynylsulfinyl, C₃-C₁₂alkynylsulfonyl, C₁-C₄alkoxycarbonyl-C₁-C₄alkylthio, C₁-C₄alkoxycarbonyl-C₁-C₄alkylsulfinyl, C_1 - C_4 alkoxycarbonyl- C_1 - C_4 alkylsulfonyl, $(C_1$ - C_4 alkoxy) $_2$ P(O)O, C_1 - C_4 alkyl- $(C_1-C_4alkoxy)P(O)O, H(C_1-C_4alkoxy)P(O)O, R_{37}R_{38}N, R_{39}R_{40}NNH, R_{41}R_{42}NC(O)O_{-1}$ R₄₃R₄₄NC(O)NH-, C₁-C₁₈alkylcarbonyloxy, C₂-C₁₈alkenylcarbonyloxy, C₂-C₁₈alkynylcarbonyloxy, C₃-C₆cycloalkylcarbonyloxy, C₁-C₁₂alkoxycarbonyloxy, C₁-C₁₂alkylthiocarbonyloxy or C₁-C₁₂alkylthiocarbamoyl, where the alkyl, alkenyl and alkynyl groups may be substituted by halogen, C_1 - C_6 alkoxy, C_1 - C_6 alkylsulfinyl, C_1 - C_6 alkylsulfonyl or cyano; or

 R_{36} is phenoxy, phenylthio, phenylsulfinyl, phenylsulfonyl, phenylsulfonylamino, phenylsulfonyloxy or benzoyloxy, where the phenyl groups for their part may be mono- or polysubstituted by halogen, nitro, cyano, C_1 - C_4 alkyl, C_1 - C_4 haloalkyl, C_1 - C_4 alkoxy, C_1 - C_4 haloalkoxy,

or R_{36} is a group Het₇-thio, Het₈-sulfinyl, Het₉-sulfonyl, Het₁₀-(CO)O or Het₁₁-N(R_{47}); in which Het₇, Het₈, Het₉, Het₁₀ and Het₁₁ independently of one another are a five- to ten-membered monocyclic or fused bicyclic ring system which may be aromatic or partially saturated and may contain 1 to 4 heteroatoms selected from the group consisting of nitrogen, oxygen and sulfur, and where each ring system may not contain more than 2 oxygen atoms and not more than 2 sulfur atoms, and where the ring system for its part may be substituted by C_1 - C_6 alkyl, C_1 - C_6 haloalkyl, C_1 - C_6 alkoxy, C_1 - C_6 haloalkoxy, C_1 - C_6 alkylsulfinyl, C_1 - C_6 alkylsulfonyl, di(C_1 - C_4 alkyl)aminosulfonyl, di(C_1 - C_4 alkyl)amino, halogen, cyano, nitro or phenyl, and where the substituents on the nitrogen in the heterocyclic ring are different from halogen;

 R_{37} , R_{38} , R_{39} , R_{40} , R_{41} , R_{42} , R_{43} , R_{44} and R_{47} independently of one another are hydrogen or C_1 - C_6 alkyl; or

 R_{37} and R_{38} together or R_{39} and R_{40} together or R_{41} and R_{42} together or R_{43} and R_{44} together are pyrolidino, piperidino, morpholino, thiomorpholino, which may be mono- or polysubstituted by methyl groups; or Q is Q_3

$$R_{50}$$
-S(O)_n R_{49} (Q₃);

in which

 R_{49} is C_1 - C_4 alkyl, C_1 - C_4 haloalkyl, C_3 - C_6 cycloalkyl or halogen-substituted C_3 - C_6 cycloalkyl; R_{50} is C_1 - C_3 alkylene which may be substituted by halogen, hydroxyl, C_1 - C_6 alkoxy, C_2 - C_6 alkenyl, C_2 - C_6 alkynyl, C_3 - C_6 cycloalkyl, C_1 - C_6 alkoxy- C_1 - C_6 alkoxy, C_1 - C_6 alkoxy- C_1 - C_6 alkoxy, (3-oxetanyl)oxy, or by C_1 - C_6 alkyl-substituted (3-oxetanyl)oxy, or by benzylthio, benzylsulfinyl, benzylsulfonyl, phenyl, phenoxy, phenylthio, phenylsulfinyl or phenylsulfonyl, where the phenyl- and benzyl-containing groups for their part may be substituted by one or more C_1 - C_6 alkyl, C_1 - C_6 haloalkyl, C_1 - C_6 alkoxy, C_1 - C_6 haloalkoxy, halogen, cyano, hydroxyl or nitro groups;

or R_{50} is phenyl, where the phenyl-containing group for its part may be substituted by one or more C_1 - C_6 alkyl, C_1 - C_6 haloalkyl, C_1 - C_6 alkoxy, C_1 - C_6 haloalkoxy, halogen, cyano, hydroxyl or nitro,

or R_{50} is C_3 - C_6 cycloalkyl, C_1 - C_6 alkoxy- or C_1 - C_6 alkyl-substituted C_3 - C_6 cycloalkyl, 3-oxetanyl or C_1 - C_6 alkyl-substituted 3-oxetanyl; and

n is 0, 1 or 2; and agronomically acceptable salts/N-oxides/isomers/enantiomers of these compounds.

The alkyl groups occurring in the definitions of the substituents can be straight-chain or branched and are, for example, methyl, ethyl, n-propyl, isopropyl, n-butyl, sec-butyl, isobutyl, tert-butyl, pentyl, hexyl, heptyl, octyl, nonyl, decyl, undecyl and dodecyl and their branched isomers. Alkoxy, alkenyl and alkynyl radicals are derived from the alkyl radicals mentioned. The alkenyl and alkynyl groups can be mono- or polyunsaturated.

Halogen is generally fluorine, chlorine, bromine or iodine, preferably fluorine and chlorine. This also applies, correspondingly, to halogen in combination with other meanings, such as haloalkyl or halophenyl.

Haloalkyl groups preferably have a chain length of from 1 to 6 carbon atoms. Haloalkyl is, for example, fluoromethyl, difluoromethyl, trifluoromethyl, chloromethyl, dichloromethyl, trichloromethyl, 2,2,2-trifluoroethyl, 2-fluoroethyl, 2-chloroethyl, pentafluoroethyl, 1,1-difluoro-2,2,2-trichloroethyl, 2,2,3,3-tetrafluoroethyl and 2,2,2-trichloroethyl; preferably trichloromethyl, difluorochloromethyl, difluoromethyl, trifluoromethyl and dichlorofluoromethyl.

Suitable haloalkenyl groups are alkenyl groups which are mono- or polysubstituted by halogen, halogen being fluorine, chlorine, bromine and iodine and in particular fluorine and chlorine, for example 2,2-difluoro-1-methylvinyl, 3-fluoropropenyl, 3-chloropropenyl, 3-chloropropenyl, 3-bromopropenyl, 2,3,3-trifluoropropenyl, 2,3,3-trichloropropenyl and 4,4,4-trifluorobut-2-en-1-yl. Among the C₃-C₂₀alkenyl groups which are mono-, di- or trisubstituted by halogen, preference is given to those having a chain length of from 3 to 5 carbon atoms.

Suitable haloalkynyl groups are, for example, alkynyl groups which are mono- or polysubstituted by halogen, halogen being bromine, iodine and in particular fluorine and chlorine, for example 3-fluoropropynyl, 3-chloropropynyl, 3-bromopropynyl, 3,3,3-trifluoropropynyl and 4,4,4-trifluorobut-2-yn-1-yl. Among the alkynyl groups which are mono- or polysubstituted by halogen, preference is given to those having a chain length of from 3 to 5 carbon atoms.

In the context of the present invention, the alkali metal cation M^{+} (for example in the definition of R_{13}) is preferably the sodium cation or the potassium cation.

Alkoxy groups preferably have a chain length of from 1 to 6 carbon atoms. Alkoxy is, for example, methoxy, ethoxy, propoxy, i-propoxy, n-butoxy, isobutoxy, sec-butoxy and tert-butoxy and also the isomeric pentyloxy and hexyloxy radicals; preferably methoxy and ethoxy. Alkylcarbonyl is preferably acetyl or propionyl. Alkoxycarbonyl is, for example, methoxycarbonyl, ethoxycarbonyl, propoxycarbonyl, isopropoxycarbonyl, n-butoxycarbonyl, isobutoxycarbonyl, sec-butoxycarbonyl or tert-butoxycarbonyl; preferably methoxycarbonyl or ethoxycarbonyl. Haloalkoxy groups preferably have a chain length of from 1 to 8 carbon

atoms. Haloalkoxy is, for example, fluoromethoxy, difluoromethoxy, trifluoromethoxy, 2,2,2-trifluoroethoxy, 1,1,2,2-tetrafluoroethoxy, 2-fluoroethoxy, 2-chloroethoxy, 2,2-difluoroethoxy and 2,2,2-trichloroethoxy; preferably difluoromethoxy, 2-chloroethoxy and trifluoromethoxy. Alkylthio groups preferably have a chain length of from 1 to 8 carbon atoms. Alkylthio is, for example, methylthio, ethylthio, propylthio, isopropylthio, n-butylthio, isobutylthio, sec-butylthio or tert-butylthio, preferably methylthio and ethylthio. Alkylsulfinyl is, for example, methylsulfinyl, ethylsulfinyl, propylsulfinyl, isopropylsulfinyl, n-butylsulfinyl, isobutylsulfinyl, sec-butylsulfinyl, tert-butylsulfinyl; preferably methylsulfinyl and ethylsulfinyl.

Alkylsulfonyl is, for example, methylsulfonyl, ethylsulfonyl, propylsulfonyl, isopropylsulfonyl, n-butylsulfonyl, isobutylsulfonyl, sec-butylsulfonyl or tert-butylsulfonyl; preferably methylsulfonyl or ethylsulfonyl. Alkoxyalkoxy groups preferably have a chain length of from 1 to 8 carbon atoms. Examples of alkoxyalkoxy groups are: methoxymethoxy, methoxyethoxy, methoxypropoxy, ethoxymethoxy, ethoxyethoxy, propoxymethoxy or butoxybutoxy. Alkylamino is, for example, methylamino, ethylamino, n-propylamino, isopropylamino or the isomeric butvlamines. Dialkylamino is, for example, dimethylamino, methylethylamino, diethylamino, n-propylmethylamino, dibutylamino and diisopropylamino. Preference is given to alkylamino groups having a chain length of from 1 to 4 carbon atoms. Alkoxyalkyl groups preferably have a chain length of 1 to 6 carbon atoms. Alkoxyalkyl is, for example, methoxymethyl, methoxyethyl, ethoxymethyl, ethoxyethyl, n-propoxymethyl, n-propoxyethyl, isopropoxymethyl or isopropoxyethyl. Alkylthioalkyl groups preferably have from 1 to 8 carbon atoms. Alkylthioalkyl is, for example, methylthiomethyl, methylthioethyl, ethylthiomethyl, ethylthioethyl, n-propylthiomethyl, n-propylthioethyl, isopropylthiomethyl, isopropylthioethyl, butylthiomethyl, butylthioethyl or butylthiobutyl. The cycloalkyl groups preferably have from 3 to 8 ring carbon atoms, for example cyclopropyl, cyclobutyl, cyclopentyl, cyclohexyl, cycloheptyl and cyclooctyl. Phenyl, also as part of a substituent such as phenoxy, benzyl, benzyloxy, benzoyl, phenylthio, phenylalkyl, phenoxyalkyl, may be substituted. In this case, the substituents can be in ortho, meta and/or para position. The preferred substituent positions are the ortho and para positions to the ring attachment point.

The compounds of the formula I may occur in different tautomeric forms, for example, if R_{13} is hydroxyl, in the preferred formulae I' and I'''

$$R_3$$
 N
 R_1
 R_2
 R_3
 R_1
 R_2
 R_3
 R_1
 R_2
 R_3
 R_2

The invention also embraces the salts which can be formed by compounds of the formula I, preferably with amines, alkali metal and alkaline earth metal bases or quarternary ammonium bases. Suitable salt formers are described, for example, in WO 98/41089.

The invention also embraces the salts which can be formed by the compounds of the formula I with amines, alkali metal and alkaline earth metal bases or quarternary ammonium bases. Among the alkali metal and alkaline earth metal hydroxides, the hydroxides of lithium, sodium, potassium, magnesium or calcium, in particular those of sodium or potassium, may be emphasized as salt formers.

Examples of amines suitable for ammonium salt formation are both ammonia and primary, secondary and tertiary C_1 - C_1 8alkylamines, C_1 - C_4 4hydroxyalkylamines and C_2 - C_4 alkoxyalkylamines, for example methylamine, ethylamine, n-propylamine, isopropylamine, the four isomeric butylamines, n-amylamine, isoamylamine, hexylamine, heptylamine, octylamine,

nonylamine, decylamine, pentadecylamine, hexadecylamine, heptadecylamine, octadecylamine, methylethylamine, methylisopropylamine, methylhexylamine, methylnonylamine, methylpentadecylamine, methyloctadecylamine, ethylbutylamine, ethylheptylamine, ethyloctylamine, hexylheptylamine, hexyloctylamine, dimethylamine, diethylamine, di-n-propylamine, disopropylamine, di-n-butylamine, di-n-amylamine, disoamylamine, dihexylamine, dibetylamine, dioctylamine, ethanolamine, n-propanolamine, isopropanolamine, N,N-diethanolamine, N-ethylpropanolamine, N-butylethanolamine, allylamine, n-butenyl-2-amine, n-pentenyl-2-amine, 2,3-dimethylbutenyl-2-amine, dibutenyl-2-amine, n-hexenyl-2-amine, propylenediamine, trimethylamine, triethylamine, tri-n-propylamine, triisopropylamine, tri-n-butylamine, triisobutylamine, tri-sec-butylamine, tri-n-amylamine, methoxyethylamine and ethoxyethylamine; heterocyclic amines, for example pyridine, quinoline, isoquinoline, morpholine, piperidine, pyrrolidine, indoline, quinuclidine and azepine; primary arylamines, for example anilines, methoxyanilines, ethoxyanilines, o,m,p-chloroanilines; but in particular triethylamine, isopropylamine and diisopropylamine.

Preferred quarternary ammonium bases which are suitable for salt formation correspond, for example, to the formula $[N(R_aR_bR_cR_d)]OH$, in which R_a , R_b , R_c and R_d independently of one another are C_1 - C_4 alkyl. Other suitable tetraalkylammonium bases with other anions can be obtained, for example, by anion exchange reactions.

Compounds of formula I, wherein p is 0, are preferred.

Preferred compounds of the formula I are those in which R₁ is -CH₂-, -CH₂CH₂-, -CF₂, -CH=CHCH₂-, -CH(CH₃)- or -C≡CCH₂-, but particularly preferably -CH₂- where in each case the free valences on the left are attached to the pyridine ring.

Preference is furthermore given to those compounds of the formula I, in which X_1 is oxygen, sulfonyl or a group $-NR_{52}SO_{2}$ -, in particular oxygen.

Of particular interest are compounds of the formula I, in which R₂ is -CH₂OCH₃, -CH₂OCH₂CH₂OCH₃, -CH₂CH₂OCH₃, -CH₂CH₂OCH₃, or -CH₂CH₂OCH₂CH₂OCH₃, preferably -CH₂CH₂OCH₃, those compounds standing out in which X₁ is oxygen and R₁ is -CH₂-.

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Among this group, preference is given to those compounds in which Q is Q_1 and R_{13} is hydroxyl.

Emphasis is furthermore given to the compounds of the formula I in which R₂

 O_{CH_2} och O_{CH_3} . If, in these preferred meanings of R_2 , no free valency is indicated, as, for example, in the case of O_{CH_3} , the point of attachment is the carbon atom indicated by "CH".

In a further preferred group of compounds of the formula I, R₃ is CF₃, CF₂CF₃, CF₂CI, CF₂H or CCI₃, particularly preferably CF₃, where R₄ is preferably hydrogen or methyl, particularly preferably hydrogen.

 R_6 , R_7 , R_8 , R_9 , R_{10} , R_{11} , R_{12} , R_{51} and R_{52} independently of one another are in particular hydrogen, C_1 - C_4 alkyl, C_1 - C_6 alkoxy- C_1 - C_6 alkyl or C_1 - C_6 alkoxy- C_1 - C_6 alkoxy, where in a preferred group of compounds of the formula I additionally Q is Q_2 and Q_3 is methylene.

Very particularly preferably, Q is Q_1 and R_{13} is hydroxyl or halogen, in particular hydroxyl. Among this group, emphasis is given to those compounds in which

- a) A_1 is $C(R_{14}R_{15})$ or NR_{16} and A_2 is $C(R_{17}R_{18})$, C(O) or oxygen, or
- b) A_1 is $C(R_{14}R_{15})$ and A_2 is $C(R_{17}R_{18})$ and R_{14} and R_{22} together form a C_2 - C_3 alkylene chain, preferably an ethylene chain, where R_{15} , R_{17} , R_{18} and R_{21} are particularly preferably hydrogen; or
- c) A_2 is C(O) or C($R_{17}R_{18}$), A_1 is C($R_{14}R_{15}$) and R_{14} , R_{15} , R_{17} and R_{18} independently of one another are hydrogen, methyl, ethyl, methoxycarbonyl or ethoxycarbonyl; or
- d) R_{14} and R_{15} or R_{21} and R_{22} together form a C_2 alkylene chain (cyclopropyl ring), A_2 is CH_2 and R_{21} and R_{22} or R_{14} and R_{15} independently of one another are hydrogen, C_1 - C_4 alkyl, methoxycarbonyl or ethoxycarbonyl; or
- e) A_2 is $C(R_{17}R_{18})$ and A_1 is $C(R_{14}R_{15})$ and R_{18} and R_{14} together form a C_2 - C_3 alkylene chain.

In a further outstanding group of compounds of the formula I, Q is Q_3 , R_{49} is cyclopropyl and R_{50} -S(O)_n is methylthio, ethylthio, methylsulfinyl, ethylsulfinyl, methylsulfonyl or ethylsulfonyl.

The compounds of the formula I can be prepared by processes known per se, for example those described in WO 97/46530 or WO 00/15615 or WO/0039094, for example in the case of compounds of the formula I,

$$\begin{array}{c|c} R_4 & O \\ \hline \\ R_3 & N & R_1 \\ \hline \\ X_1 & R_2 \end{array} \qquad (I)$$

in which $R_1,\,R_2,\,R_3,\,R_4$ and X_1 are as defined under formula I and Q is a group

$$\begin{array}{c} (Q_1), \\ R_{22} \\ A_2 \\ A_1 \end{array} O$$

by, for example, either

a) reacting a compound of the formula la

$$R_4$$
 N
 R_1
 X_1
 R_2
(la),

in which R_1 , R_2 , R_3 , R_4 and X_1 are as defined under formula I and Y_1 is a leaving group, for example halogen or cyano, in an inert organic solvent in the presence of a base with a compound of the formula II

in which R_{22} , R_{21} , A_2 and A_1 are as defined under formula I, to give the compounds of the formulae IIIa and IIIb

and then isomerizing these for example in the presence of a base and a catalytic amount of dimethylaminopyridine (DMAP) or a source of cyanide, for example acetone cyanohydrin; or b) reacting a compound of the formula lb

$$R_4$$
 OH R_3 N R_1 X_1 R_2 $(Ib),$

in which R_1 , R_2 , R_3 , R_4 and X_1 are as defined under formula I, with a compound of the formula II

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in which R_{22} , R_{21} , A_1 and A_2 are as defined under formula I, in an inert organic solvent in the presence of a base and a coupling agent to give the compounds of the formula IIIa or IIIb

$$R_4$$
 R_2
 R_3
 R_1
 R_2
 R_3
 R_4
 R_2
 R_3
 R_4
 R_3
 R_4
 R_5
 R_5

and then isomerizing these, for example as described under route a).

Compounds of the formula I, in which Q is a group

are prepared similarly to a known process (for example WO 97/46530), wherein either

a) a compound of the formula la

$$R_4$$
 R_3
 N
 R_1
 X_1
 R_2
(la),

in which R_1 , R_2 , R_3 , R_4 and X_1 are as defined under formula I and Y_1 is a leaving group, for example halogen or cyano, is reacted with a compound of the formula IIa

$$R_{34}$$
 N
 N
 O
(IIa),
 R_{35}

in which R_{34} and R_{35} are as defined, in an inert organic solvent in the presence of a base to give the compound of the formula IIIc

$$R_4$$
 R_3
 R_3
 R_3
 R_3
 R_3
 R_3
 R_2
IIIc

in which R_1 , R_2 , R_3 , R_4 , R_{34} , R_{35} and X_1 are as defined under formula I, and this compound is then isomerized, for example in the presence of a base and a catalytic amount of a source of cyanide; or

b) a compound of the formula lb

$$R_4$$
 OH R_3 N R_1 R_2 (lb),

in which R_1 , R_2 , R_3 , R_4 and X_1 are as defined under formula I, is reacted with a compound of the formula IIa

in which R_{34} and R_{35} are as defined above, in an inert organic solvent in the presence of a base and a coupling agent to give the compound of the formula IIIc

$$R_{3}$$
 R_{3}
 R_{3}

and this compound is then isomerized as described under route a).

The compounds of the formula I, in which Q is a group

in which n is 0 and R_{50} and R_{49} are as defined above, are prepared similarly to known processes (for example those described in WO 00/15615, WO/0039094 or WO 97/43270), wherein either

a) a compound of the formula IV

$$\begin{array}{c|c} R_{4} & & \\ R_{3} & N & R_{1} \\ & & \\ X_{1} & & \\ & & R_{2} \end{array} \qquad \text{(IV),}$$

in which X_1 , R_1 , R_2 , R_3 , R_4 and R_{49} are as defined above, is converted in the presence of a base, carbon disulfide and an alkylating agent of the formula V

$$R_{50}$$
- Y_2 (V),

in which R_{50} is as defined under formula I, and Y_2 is a leaving group, for example halogen or sulfonate, into the compound of the formula VI

$$\begin{array}{c|c} & & & & \\ & & & & \\ & & & & \\ R_3 & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ &$$

in which R₁, R₂, R₃, R₄, R₅₀, X₁ and R₄₉ are as defined above, and this compound is then cyclized with hydroxylamine hydrochloride, if appropriate in a solvent, in the presence of a base, for example sodium acetate, to give the isomeric compounds of the formulae Ic and Id

$$\begin{array}{c} (O)n \\ S \\ R_{50} \\ O \\ R_{49} \\ R_{3} \\ N \\ R_{1} \\ R_{2} \\ \end{array} \qquad \text{and} \qquad \begin{array}{c} (O)n \\ S \\ R_{49} \\ R_{49} \\ R_{3} \\ \end{array} \qquad \begin{array}{c} (O)n \\ S \\ R_{50} \\ O \\ R_{49} \\ R_{3} \\ \end{array} \qquad \begin{array}{c} (O)n \\ S \\ R_{49} \\ R_{49} \\ R_{1} \\ X_{1} \\ R_{2} \\ \end{array} \qquad \qquad \begin{array}{c} (Ic) \\ (Id) \\ \end{array} \qquad \begin{array}{c} (O)n \\ S \\ R_{49} \\ R_{49} \\ R_{1} \\ X_{1} \\ R_{2} \\ \end{array} \qquad \begin{array}{c} (O)n \\ S \\ R_{49} \\ R_{1} \\ X_{1} \\ R_{2} \\ \end{array} \qquad \begin{array}{c} (O)n \\ S \\ R_{49} \\ R_{1} \\ X_{1} \\ R_{2} \\ \end{array} \qquad \begin{array}{c} (Ic) \\ R_{1} \\ R_{2} \\ \end{array} \qquad \begin{array}{c} (Ic) \\ R_{2} \\ R_{2} \\ \end{array} \qquad \begin{array}{c} (Id) \\ R_{2} \\ \end{array} \qquad \begin{array}{c} (Id) \\ R_{2} \\ R_{2} \\ \end{array} \qquad \begin{array}{c} (Id) \\ R_{2} \\ R_{3} \\ \end{array} \qquad \begin{array}{c} (Id) \\ R_{2} \\ R_{3} \\ \end{array} \qquad \begin{array}{c} (Id) \\ R_{2} \\ R_{3} \\ \end{array} \qquad \begin{array}{c} (Id) \\ R_{3} \\ R_{4} \\ R_{3} \\ \end{array} \qquad \begin{array}{c} (Id) \\ R_{4} \\ R_{3} \\ R_{4} \\ \end{array} \qquad \begin{array}{c} (Id) \\ R_{4} \\ R_{4} \\ R_{5} \\ R_$$

and these compounds are then oxidized with an oxidizing agent, for example with peracids, for example meta-chloroperbenzoic acid (m-CPBA) or peracetic acid, to give the corresponding sulfoxides (n = 1) and sulfones (n = 2) of the formulae le and lf, respectively. Isomers of the formulae lc and ld (in which n = 0) or le and lf (in which n = 1 or 2) can be separated and purified by column chromatography, using a suitable mobile phase.

The intermediates of the formulae Ia, Ib, IV and VI are novel and were developed specifically for the preparation of the compounds of the formula I. Accordingly, they also form part of the subject-matter of the present invention. Together, the novel intermediates of the formulae Ia, Ib, IV and VI correspond to formula XX

in which

Q is hydroxyl, halogen, cyano or C₁-C₆alkoxy, or is a group of the formula

$$R_{49}$$
 or $-CH_2(CO)R_{49}$; and R_{50}

R₁, R₃, R₄, R₄₉, R₅₀, X₁ and p are as defined under formula I and R₂ is a C₁-C₈alkyl. C₃-C₆alkenyl or C₃-C₆alkynyl group which is mono- or polysubstituted by halogen, hydroxyl, amino, formyl, nitro, cyano, mercapto, carbamoyl, C₁-C₆alkoxy, C₁-C₆alkoxycarbonyl C2-C6alkenyl, C2-C6haloalkenyl, C2-C6alkynyl, C2-C6haloalkynyl, C3-C6cycloalkyl, by halogensubstituted C₃-C₆cycloalkyl, or by C₃-C₆alkenyloxy, C₃-C₆alkynyloxy, C₁-C₆haloalkoxy, C₃-C₆haloalkenyloxy, cyano-C₁-C₆alkoxy, C₁-C₆alkoxy-C₁-C₆alkox C_1 - C_6 alkoxy, C_1 - C_6 a C_1 - C_6 alkoxy, C_1 - C_6 alkoxycarbonyl- C_1 - C_6 alkoxy, C_1 - C_6 alkoxycarbonyl, C_1 - C_6 alkoxycarbonyl, C_1 - C_6 alkylthio, C_1 - C_6 alkylsulfinyl, C_1 - C_6 alkylsulfonyl, C_1 - C_6 haloalkylthio, C₁-C₆haloalkylsulfinyl, C₁-C₆-haloalkylsulfonyl, oxiranyl, which for its part may be substituted by C₁-C₆alkyl, or by (3-oxetanyl)oxy, which for its part may be substituted by C₁-C₆alkyl, or by benzylthio, benzylsulfinyl, benzylsulfonyl, C₁-C₆alkylamino, di(C₁-C₆alkyl)amino, R₉S(O)₂O, $R_{10}N(R_{11})SO_2$ -, thiocyanato, phenyl, phenoxy, phenylthio, phenylsulfinyl or phenylsulfonyl: where the phenyl- or benzyl-containing groups for their part may be substituted by one or more C₁-C₆alkyl, C₁-C₆haloalkyl, C₁-C₆haloalkoxy, halogen, cyano, hydroxyl or nitro groups, or

 R_2 is phenyl which may be mono- or polysubstituted by C_1 - C_6 alkyl, C_1 - C_6 haloalkyl, C_1 - C_6 alkoxy, C_1 - C_6 haloalkoxy, halogen, cyano, hydroxyl or nitro; or R_2 is C_3 - C_6 cycloalkyl, C_1 - C_6 alkoxy- or C_1 - C_6 alkyl-substituted C_3 - C_6 cycloalkyl, 3-oxetanyl or C_1 - C_6 alkyl-substituted 3-oxetanyl; or

if X_1 is -N(R₆)-O-, -O-NR₅₁, SO₂NR₇- or -NR₅₂SO₂- and R₆, R₇, R₅₁ and R₅₂ are as defined under formula I,

 R_2 may additionally be hydrogen, unsubstituted C_1 - C_6 alkyl, or

a five- to ten-membered monocyclic or fused bicyclic ring system which may be aromatic, saturated or partially saturated and may contain 1 to 4 heteroatoms selected from the group consisting of nitrogen, oxygen and sulfur, where the ring system is attached directly or via a C_1 - C_4 alkylene, C_2 - C_4 alkenyl- C_1 - C_4 alkylene, C_2 - C_4 alkylene, C_2 - C_4 alkylene, C_2 - C_4 alkylene group to the substituent X_1 , and where each ring system may not contain more than 2 oxygen atoms and not more than two sulfur atoms, and where the ring system for its part may be mono-, di- or trisubstituted by C_1 - C_6 alkyl, C_1 - C_6 haloalkyl, C_2 - C_6 alkenyl, C_2 - C_6 haloalkynyl, C_3 - C_6 alkoxy, hydroxyl, C_1 - C_6 haloalkoxy, C_3 - C_6 alkenyloxy, C_3 - $C_$

 C_3 - C_6 alkynylthio, C_2 - C_5 alkoxyalkylthio, C_3 - C_5 acetylalkylthio, C_3 - C_6 alkoxycarbonylalkylthio, C_2 - C_4 cyanoalkylthio, C_1 - C_6 alkylsulfinyl, C_1 - C_6 haloalkylsulfinyl, C_1 - C_6 alkylsulfonyl, aminosulfonyl, C_1 - C_6 haloalkylsulfonyl, aminosulfonyl, C_1 - C_2 alkylaminosulfonyl, di(C_1 - C_2 alkyl)amino, halogen, cyano, nitro, phenyl and benzylthio, where phenyl and benzylthio for their part may be substituted on the phenyl ring by C_1 - C_3 alkyl, C_1 - C_3 haloalkyl, C_1 - C_3 alkoxy, C_1 - C_3 haloalkoxy, halogen, cyano or nitro, and where the substituents on the nitrogen in the heterocyclic ring are different from halogen.

The preparation of the compounds of the formula I is illustrated in more detail in reaction schemes 1 and 2 below.

Reaction scheme 1

Route b):

base e.g.
$$(C_2H_5)_3N$$
, base e.g. $(C_2H_5)_3N$, coupling agent e.g. III or $IIII$ or $IIII$ II

Reaction scheme 1 is preferably used to prepare the compounds of the formula I having the group Q_1 , in which R_{13} is hydroxyl, and the compounds of the formula I having the group Q_2 , in which R_{36} is hydroxyl.

Reaction scheme 2

$$\begin{array}{c} R_4 \\ R_3 \\ N \\ R_1 \\ N \\ R_2 \\ N \\ R_3 \\ N \\ R_1 \\ N \\ R_2 \\ N \\ N_1 \\ N_2 \\ N_1 \\ N_2 \\ N_2 \\ N_3 \\ N_4 \\ N_1 \\ N_2 \\ N_2 \\ N_3 \\ N_4 \\ N_1 \\ N_2 \\ N_4 \\ N_2 \\ N_4 \\ N_5 \\ N_5 \\ N_5 \\ N_6 \\$$

Compounds of the formula I, in which p is 1, i.e. the corresponding N-oxides of the formula I, can be prepared by reacting a compound of the formula I, in which p is 0, with a suitable oxidizing agent, for example the H_2O_2 /urea adduct in the presence of an acid anhydride, e.g. trifluoroacetic anhydride. Such oxidations are known from the literature, for example from *J. Med. Chem.*, 32 (12), 2561-73, 1989 or WO 00/15615.

According to reaction scheme 1, route a), the carboxylic acid derivatives of the formula Ia in which Y_1 is a leaving group such as halogen, for example iodine, bromine, and in particular chlorine, N-oxyphthalimide or N,O-dimethylhydroxylamino or part of an activated ester, for example N = C = N + C (formed from dicyclohexylcarbodiimide (DCC) and the

corresponding carboxylic acid) or $C_2H_5N=C-NH(CH_2)_3N(CH_3)_2$ (formed from N-ethyl N'-(3-0-

dimethylaminopropyl)carbodiimide (EDC) and the corresponding carboxylic acid) are used as starting materials for preparing the compounds of the formula I in which Q denotes the groups Q_1 and Q_2 and R_{13} and R_{36} are hydroxyl. The starting materials are reacted in an inert organic solvent such as a halogenated hydrocarbon, for example dichloromethane, a nitrile, for example acetonitrile, or an aromatic hydrocarbon, for example toluene, and in the presence of a base such as an alkylamine, for example triethylamine, an aromatic amine, for example pyridine or 4-dimethylaminopyridine (DMAP) with the dione derivatives of the formula II or pyrazoles of the formula IIa, to give the isomeric enol ethers of the formula IIIa, IIIb or IIIc. This esterification can be carried out at temperatures of from 0°C to 110°C.

The isomerization of the ester derivatives of the formulae IIIa, IIIb and IIIc to derivatives of the formula I (in which R_{13} and R_{36} are hydroxyl) can be carried out, for example, similarly to EP-A-0 353 187, EP-A-0 316 491 or WO 97/46530 in the presence of a base such as an alkylamine, for example triethylamine, a carbonate, for example potassium carbonate, and a catalytic amount of DMAP or a source of cyanide, such as acetone cyanohydrin or potassium cyanide. In particular if a cyanide compound of the formula Ia ($Y_1 = \text{cyano}$) is used, or in the presence of a catalytic amount of acetone cyanohydrin or potassium cyanide, the two reaction steps can be carried out *in situ* without isolating the intermediates III.

According to reaction scheme 1, route b), the desired derivatives of the formula I (in which R_{13} and R_{36} are hydroxyl) can be obtained, for example, similarly to E. Haslem, *Tetrahedron*, 2409-2433, *36*, 1980, by esterifying the carboxylic acids of the formula Ib with the dione derivatives of the formula II or pyrazoles of the formula IIa in an inert solvent such as a halogenated hydrocarbon, for example dichloromethane, a nitrile, for example acetonitrile, or an aromatic hydrocarbon, for example toluene, in the presence of a base such as an alkylamine, for example triethylamine, and a coupling agent such as 2-chloro-1-methyl-pyridinium iodide. Depending on the solvent used, this esterification is carried out at temperatures of from 0°C to 110°C, giving initially, as described under route a), the isomeric ester of the formula IIIa, IIIb or IIIc, which can be isomerized as described under route a), for example in the presence of a base and a catalytic amount of DMAP, or a source of cyanide, for example acetone cyanohydrin, to give the desired derivative of the formula I (R_{13} and R_{36} = hydroxyl). The activated carboxylic acid derivatives of the formula Ia in reaction scheme 1 (route a), in which Y_1 is a leaving group such as halogen, for example bromine, iodine or, in particular, chlorine, can be prepared by known standard processes, for example

those described in C. Ferri "Reaktionen der organischen Synthese" [Reactions of organic sythesis], Georg Thieme Verlag, Stuttgart, 1978, page 460 ff. Such reactions are generally known and described in the literature in different variations with respect to the leaving group Y₁.

The preparation of the compounds of the formula I, in which Q denotes the group Q_3 , can be carried out according to reaction scheme 2 by reacting the β -diketone derivative of the formula IV for example similarly to *Synthesis* 1991, 301; ibid. 1988, 793; or *Tetrahedron 32*, 3055, 1976, with carbon disulfide in the presence of a base such as a carbonate, for example potassium carbonate, a metal hydride, for example sodium hydride, or potassium fluoride on aluminum, and an alkylating agent of the formula V, in which Y_2 is a leaving group such as halogen, for example iodine, bromine and, in particular, chlorine, $CH_3SO_2O_7$ or

$$\mathrm{CH_3}$$
 $\mathrm{SO_2O}$. This reaction is expediently carried out in a solvent such as an

amide, for example N,N-dimethylformamide (DMF), a sulfoxide, for example dimethyl sulfoxide (DMSO), or a nitrile, for example acetonitrile. The ketene thioacetal of the formula VI that is formed is cyclized with the aid of hydroxylamine hydrochloride in the presence of a base such as sodium acetate in a solvent such as an alcohol, for example ethanol, or an ether, for example tetrahydrofuran, to give the isomeric compounds of the formulae Ic and Id (in which n is 0). This cyclization reaction is carried out at temperatures of from 0°C to 100°C. If appropriate, the compounds of the formulae Ic and Id in which n is 0 can be oxidized similarly to known standard processes as described, for example, in H. O. House, "Modern Synthetic Reactions", W. A. Benjamin, Inc., Menlo Park, California, 1972, pages 334-335 and 353-354, to give the corresponding sulfones and sulfoxides of the formulae Ie and If (n = 1 or 2).

The compounds of the formula IV in reaction scheme 2 can be obtained by standard processes for example from the corresponding compounds of the formula la

$$R_4$$
 Y_1
 R_3
 N
 R_1
 X_1
 R_2
(la),

in which R_1 , R_2 , R_3 , R_4 , and X_1 are as defined above and Y_1 is a leaving group, for example halogen, for example by Claisen condensation, or from the compounds of the formula la by reaction with a ketocarboxylic acid salt of the formula VII

in which R_{49} is as defined under formula I and M^{+} is an alkali metal ion (cf., for example, WO 96/26192).

Compounds of the formula I, in which R₁ is, in particular C₁-C₂alkyl, can, for example, also be prepared by heating an N-oxide of the formula IX under known reaction conditions in the presence of an acid anhydride (see, for example, Konno, K.; Hashimoto, K.; Shirahama, H.; Matsumoto, T.; Heterocycles 1986, 24, 2169, or WO 00/15615) and hydrolyzing the resulting products (Ig) in a protic solvent, for example water or a water/methanol mixture, if appropriate in the presence of a base (for example lithium hydroxide or sodium hydroxide), and then converting the resulting alcohol X in the presence of a base, for example sodium hydride or potassium hydroxide, if appropriate in the presence of a phase-transfer catalyst or a crown ether, and an alkylating agent R_2 - Y_3 , in which R_2 is as defined under formula I and Y₃ is a leaving group, for example halogen or methyl sulfonate, in an aprotic solvent, for example, tetrahydrofuran or dimethylformamide, into the corresponding derivatives of the formula Ih (in which X_1 is oxygen). Compounds of the formula I, in which R_2 is C₁-C₀alkoxymethyl or 2-tetrahydropyranyl or 2-tetrahydrofuryl, can be prepared, for example, by treating an alcohol of the formula X with a vinyl ether of the formula VE1, in which R03, R04, and R_{05} are C_1 - C_6 alkyl or R_{03} together with R_{05} forms a C_2 - C_3 alkylene chain, in the presence of an acidic catalyst, for example para-toluenesulphonic acid, in an inert solvent, for example methylene chloride. Such reactions are generally known in the literature (see, for example, Synthesis, p. 169, 1973). The two reaction sequences are demonstrated using the example below:

Reaction scheme 3

OH O oxidizing agent e.g.
$$H_2O_2$$
 H_2NCONH_2 R_2 $Solvent$ e.g. CH_2Cl_2 R_2 $Solvent$ R_3 R_4 R_4 R_5 R_5

Compounds of the formula I, in which R_1 is, in particular, C_1 - C_2 alkyl or C_1 - C_2 haloalkyl, can, for example, also be prepared by oxidzing a compound of the formula XI, in which R_{13} is in particular chlorine, C_1 - C_4 alkoxycarbonyloxy or benzoylcarbonyloxy (prepared similarly to WO 00/15615 or WO/0039094), under known halogenation conditions using, for example, N-bromosuccinimide or N-chlorosuccinimide in the presence of light and a free-radical initiator such as benzoyl peroxide to give the 1-bromo or 1-chloro, 1,1-dibromo or 1,1-dichloro compound and then refunctionalizing these compounds into the corresponding derivatives of the formula I, for example by reaction with a nucleophile R_2 -Z, in which Z is, for example, -SH, -OH, -C(O)OH, -O-N(R_{51})H, -N(R_{6})-OH, -SO₂N(R_{52})H or -N(R_{8})H and R_2 , R_{52} ,

 R_8 , R_6 and R_{51} are as defined under formula I, in the presence of a base, for example sodium hydride, potassium hydroxide or potassium carbonate, followed by aqueous work-up. These reaction sequences, too, are demonstrated by the example below.

Reaction scheme 4

$$\begin{array}{c} R_{4} \\ R_{3} \\ R_{01} \\ R_{01}$$

Compounds of the formula I, in which Q denotes Q_1 or Q_2 and in which R_{13} or R_{36} are different from hydroxyl or halogen, can be prepared by conversion processes generally known from the literature, for example acylations or carbamoylations with appropriate acid chlorides from compounds of the formula I, in which R_{13} or R_{36} is hydroxyl, in the presence of a suitable base, or they can be prepared by nucleophilic substitution reactions on chlorides of the formula I, in which R_{13} and R_{36} are chlorine, the chlorides likewise being obtainable according to known processes by reaction with a chlorinating agent, such as phosgene, thionyl chloride or oxalyl chloride. The starting materials used are, for example, appropriately substituted amines, or hydroxylamines directly, or alkylsulfonamides, mercaptans,

thiophenols, phenols, heterocyclic amines or heterocyclic thiols in the presence of a base, for example 5-ethyl-2-methylpyridine, diisopropylethylamine, triethylamine, sodium bicarbonate, sodium acetate or potassium carbonate.

Compounds of the formula I, in which R_{13} and R_{36} contain thio groups, can be oxidized similarly to known standard processes using, for example, peracids, for example meta-chloroperbenzoic acid (m-CPBA) or peracetic acid, to give the corresponding sulfones and sulfoxides of the formula I. The degree of oxidation at the sulfur atom (SO- or SO₂-) can be controlled by the amount of oxidizing agent.

The resulting derivatives of the formula I, in which R_{13} and R_{36} are different from hydroxyl, can likewise occur in different isomeric forms which, if appropriate, can be isolated in pure form. Accordingly, the invention also embraces all of these stereoisomeric forms. Examples of these isomeric forms are the formulae I*, I** and I*** below in which Q denotes the group Q_1 .

The compounds of the formulae II and IIa are known and can be prepared similarly as described, for example, in WO 92/07837, JP 10265441, DE-A-3818958, EP-A-0 338 992, DE-A-3902818, EP-A-0 278 742, WO 98/29412, JP 02059566, US-A-5,089,046, GB-A-2205316, WO 00/27821 or EP-A-0 384 736.

The required intermediates of the formula lb (or lk, ll or lm) are synthesized similarly to known processes as described, for example, in WO 00/15615, WO/00/39094 or WO 97/46530, or they can be prepared for example, according to generally known conversion methods such as the Stille (see, for example *Angew. Chem.* 1986, 98(6), 504-19), Heck (see, for example, *Angew. Chem.* 1994, 106 (23/24), 2473-506), Sonogashira (see, for example, "Comprehensive Organometallic Synthesis", Pergamon Verlag, Oxford, Vol 3, 1991, page 521 ff.) or Wittig (for example C. Ferri "Reaktionen der organischen Synthese", Georg Thieme Verlag, Stuttgart, 1978, p. 354 ff.) reactions, starting from halogen derivatives of the formula XIV (preparation as described in WO 00/15615 or WO/0039094) or XVII (preparation similar to EP 522392) (reaction scheme 5):

Reaction scheme 5

$$R_{0z} = C_1 - C_8 alkyl, hydrogen \\ R_{0z} =$$

Intermediates of the formula lb, in which R_1 , R_2 , R_3 , R_4 and X_1 are as defined under formula I, can also be prepared by the method according to reaction scheme 6:

Reaction scheme 6

$$\begin{array}{c} R_s - Y_3 \\ \text{base e.g. NaH} \\ \text{or KOH/crown ether, e.g. 18-C-6} \\ R_3 - N - (CH_2)n_{\infty} \\ \text{Desce e.g. NaH} \\ \text{CO}_2R_{\infty} \\ \text{R}_3 - N - (CH_2)n_{\infty} \\ \text{Desce e.g. NaH} \\ \text{Solidation} \\ \text{e.g. perbenzole acid} \\ \text{e.g. perbenzole acid} \\ \text{e.g. perbenzole acid} \\ \text{R}_3 - N - (CH_2)n_{\infty} \\ \text{R}_4 - CO_2R_{\infty} \\ \text{R}_5 - N - (CH_2)n_{\infty} \\ \text{R}_5 - N - (CH_2)n_{$$

Using generally known oxidation methods such as dihydroxylation, Wacker oxidation, epoxidation, hydroboration with subsequent oxidation, starting with vinyl or allyl compounds of the formula XVIII (preparation as described in WO 00/15615 or WO/0039094), intermediates of the formulae II, In, Iq and Ir are obtained which can be converted by conversion processes known to the person skilled in the art (for example alcohol activation, for example as sulfonate, alkylation, for example using an alkylating agent R_2 - Y_3 or R_5 - Y_3 , in which R_2 and R_5 are as defined under formula I and Y_3 is a leaving group, for example halogen), in the presence of a base, or using nucleophile reactions, for example with a nucleophile Z- R_2 , in which Z and R_2 are as defined above, into compounds of the formula I.

Intermediates of the formula Ib, in which R_1 is C_1 - C_2 alkyl and R_2 , R_3 , R_4 and X_1 are as defined under formula I, can also be prepared by reacting a compound of the formula XIVa, in which R_3 and R_4 are as defined above under formula I and Y_4 is halogen, with a nucleophile R_2 -Z, in which Z is -SH, -OH, -C(O)OH, -O-N(R_{51})H, -N(R_6)-OH -SO $_2$ N(R_{52})H or -N(R_8)H and R_2 , R_{52} , R_8 , R_6 , R_{51} are as defined above under formula I, in the presence of a base such as sodium hydride or an alkaline earth metal oxide or carbonate in an inert solvent such as dimethylformamide or THF at temperatures between -5 and 160°C, or, to prepare the corresponding sulfinyl or sulfonyl derivatives of the formula Iu, by reacting with an oxidizing agent such as m-chloroperbenzoic acid or sodium periodate, or sodium perborate, with, depending on the degree of oxidation, temperature control known to the person skilled in the art (for example -30°C-+50°C for n=1 and -20°C-+100°C for n=2 respectively), in an inert solvent such as dichloromethane, to give compound of the formula Iv. In reaction scheme 7 below, this is illustrated in more detail for the case Z = OH, SH, $SO_2N(R_{52})H$ and $N(R_8)H$:

Reaction scheme 7

Intermediates of the formula I, in which Q denotes a group OR₀₂ (R₀₂ = C₁-C₆alkyl), can be converted by hydrolysis using, for example, a base, for example LiOH, in a protic solvent, for example H₂O or H₂O/methanol mixtures, into products of the formula lb.

For preparing all further compounds of the formula I functionalized according to the definitions of R₁, R₂, R₃, R₄ and X₁, there are a large number of suitable known standard methods, for example alkylation, halogenation, acylation, amidation, oximation, oxidation and reduction, the choice of the preparation methods which are suitable depending on the properties (reactivity) of the substituents in the intermediates.

The reactions to give compounds of the formula I are advantageously carried out in aprotic inert organic solvents. Such solvents are hydrocarbons such as benzene, toluene, xylene or cyclohexane, chlorinated hydrocarbons such as dichloromethane, trichloromethane, tetrachloromethane or chlorobenzene, ethers such as diethyl ether, ethylene glycol dimethyl ether, diethylene glycol dimethyl ether, tetrahydrofuran or dioxane, nitriles such as acetonitrile or propionitrile, amides such as N,N-dimethylformamide, diethylformamide or N-methylpyrrolidinone. The reaction temperatures are advantageously between -20°C and +120°C. In general, the reactions are slightly exothermic and, as a rule, they can be carried out at room temperature. To shorten the reaction time, or else to start the reaction, the mixture may be heated briefly to the boiling point of the reaction mixture. The reaction times can also be shortened by adding a few drops of base as reaction catalyst. Suitable bases are, in particular, tertiary amines such as trimethylamine, triethylamine, quinuclidine, 1,4-diazabicyclo[2.2.2]octane, 1,5-diazabicyclo[4.3.0]non-5-ene or 1,5-diazabicyclo-[5.4.0]undec-7-ene. However, inorganic bases such as hydrides, e.g. sodium hydride or calcium hydride, hydroxides, e.g. sodium hydroxide or potassium hydroxide, carbonates such as sodium carbonate and potassium carbonate, or hydrogen carbonates such as potassium hydrogen carbonate and sodium hydrogen carbonate may also be used as bases. The bases can be used as such or else with catalytic amounts of a phase-transfer catalyst, for example a crown ether, in particular 18-crown-6, or a tetraalkylammonium salt.

The compounds of the formula I can be isolated in the customary manner by concentrating and/or by evaporating the solvent and purified by recrystallization or trituration of the solid residue in solvents in which they are not readily soluble, such as ethers, aromatic hydrocarbons or chlorinated hydrocarbons.

All application methods which are conventionally used in agriculture, for example preemergence application, post-emergence application and seed treatment, as well as various methods and techniques, for example the controlled release of active ingredients, are suitable for the use according to the invention of the compounds of the formula I or of compositions comprising them. To this end, the active ingredient in solution is applied to mineral carriers for granules or to polymerized granules (urea/formaldehyde) and dried. If appropriate, an additional coating can be applied (coated granules), which allows the active ingredient to be released in a controlled manner over a specific period of time.

The compounds of the formula I can be employed as herbicides as such, i.e. as obtained from synthesis. However, they are preferably processed in the customary manner together with the auxiliaries conventionally used in the art of formulation, for example to give emulsifiable concentrates, directly sprayable or dilutable solutions, dilute emulsions, wettable powders, soluble powders, dusts, granules or microcapsules. Such formulations are described, for example, in WO 97/34485 on pages 9 to 13. The application methods such as

spraying, atomizing, dusting, wetting, scattering or pouring, as well as the type of composition, are chosen to suit the intended aims and the prevailing circumstances.

The formulations, i.e. the compositions, preparations or products which comprise the active ingredient of the formula I or at least one active ingredient of the formula I and, as a rule, one or more solid or liquid formulation auxiliaries, are prepared in the known manner, for example by intimately mixing and/or grinding the active ingredients together with the formulation auxiliaries, for example solvents or solid carriers. Furthermore, surface-active compounds (surfactants) may additionally be used when preparing the formulations. Examples of solvents and solid carriers are indicated for example in WO 97/34485 on page 6.

Suitable surface-active compounds are, depending on the nature of the active ingredient of the formula I to be formulated, nonionic, cationic and/or anionic surfactants and surfactant mixtures which have good emulsifying, dispersing and wetting properties. Examples of suitable anionic, nonionic and cationic surfactants are enumerated, for example, in WO 97/34485 on pages 7 and 8. The surfactants conventionally used in the art of formulation which are described, inter alia, in "McCutcheon's Detergents and Emulsifiers Annual" MC Publishing Corp., Ridgewood New Jersey, 1981, Stache, H., "Tensid-Taschenbuch" ["Surfactants Guide"], Carl Hanser Verlag, Munich/Vienna, 1981 and M. and J. Ash, "Encyclopedia of Surfactants", Vol I-III, Chemical Publishing Co., New York, 1980-81 are furthermore also suitable for preparing the herbicidal compositions according to the invention.

As a rule, the herbicidal formulations comprise 0.1 to 99% by weight, in particular 0.1 to 95% by weight, of herbicide, 1 to 99.9% by weight, in particular 5 to 99.8% by weight, of a solid or liquid formulation auxiliary and 0 to 25% by weight, in particular 0.1 to 25% by weight, of a surfactant. While concentrated compositions are more preferred as commercially available goods, the end consumer uses, as a rule, dilute compositions. The compositions can also comprise further additives such as stabilizers, for example epoxidized or non-epoxidized vegetable oils (epoxidized coconut oil, rapeseed oil or soya oil), antifoams, e.g. silicone oil, preservatives, viscosity regulators, binders, tackifiers and fertilizers or other active ingredients.

As a rule, the active ingredients of the formula I are applied to the plant or its environment at rates of 0.001 to 4 kg/ha, in particular 0.005 to 2 kg/ha. The dosage required for the desired

action can be determined by experiments. It depends on the type of the action, the developmental stage of the crop plant and of the weed, and on the application (location, timing, method) and can, owing to these parameters, vary within wide limits.

The compounds of the formula I are distinguished by herbicidal and growth-inhibitory properties which allow them to be employed in crops of useful plants, in particular in cereals, cotton, soya, sugar beet, sugar cane, plantation crops, rapeseed, maize and rice and for the non-selective control of weeds. Crops are also to be understood as including those which have been rendered tolerant to herbicides or classes of herbicides by means of conventional plant-breeding or genetic-engineering methods. The weeds to be controlled may be both mono- and dicotyledonous weeds such as Stellaria, Nasturtium, Agrostis, Digitaria, Avena, Setaria, Sinapis, Lolium, Solanum, Echinochloa, Scirpus, Monochoria, Sagittaria, Bromus, Alopecurus, Sorghum halepense, Rottboellia, Cyperus, Abutilon, Sida, Xanthium, Amaranthus, Chenopodium, Ipomoea, Chrysanthemum, Galium, Viola and Veronica.

The examples which follow illustrate the invention in greater detail without limiting it.

Preparation examples:

Example H1: Preparation of ethyl 2-bromomethyl-6-trifluoromethylnicotinate:

434.4 g (1.866 mol) of ethyl 2-methyl-6-trifluoromethylnicotinate (preparation similar to Heterocycles 129, 46, 1997) and 398.5 g (2.239 mol) of N-bromosuccinimide in 3 500 ml of carbon tetrachloride in the presence of 30.6 g (0.1866 mol) of α , α -azaisobutyronitrile are heated at 75°C, with irradiation from a 150 Watt lamp. After 3 hours, the reaction is terminated, the mixture is cooled to 15°C and precipitated succinimide is removed by filtration. After evaporation of the solvent, the residue is distilled under reduced pressure. This gives ethyl 2-bromomethyl-6-trifluoromethylnicotinate as an oily product (260.2 g, 44.7% of theory, b.p. 74°C/0.04 mmHg).

Example H2: 2-(2-Methoxyethoxymethyl)-6-trifluoromethylnicotinic acid:

At room temperature,177.2 g of ethyl 2-bromomethyl-6-trifluoromethylnicotinate are dissolved in 3 000 ml of toluene and reacted with 398 ml (1.704 mol) of a 21% ethanolic solution of sodium ethoxide. After 8 hours at room temperature, 1500 ml of ethanol and 100 ml of 30% aqueous sodium hydroxide solution are added with vigorous stirring, and the

reaction mixture is stirred at this temperature for another 4 hours. The reaction mixture is poured into water and extracted with ethyl acetate, and the aqueous phase is acidified to pH 1. Following extraction with ethyl acetate, drying over sodium sulfate, evaporation under reduced pressure and trituration with hexane, pure 2-(2-methoxyethoxymethyl)-6-trifluoromethylnicotinic acid is obtained in the form of white crystals of melting point 62-63°C.

Example H3: 4-Hydroxy-3-[2-(2-methoxyethoxymethyl)-6-trifluoromethylpyridine-3-carbonyl]-bicyclo[3.2.1]oct-3-en-2-one:

24.9 g (0.1 mol) of 2-(2-methoxyethoxymethyl)-6-trifluoromethylnicotinic acid are dissolved in 200 ml of methylene chloride and 20 ml of oxalyl chloride, and 0.1 ml of dimethylformamide is then added dropwise. After the strong evolution of gas has ceased, triethylamine (27.9 ml, 0.2 mol), dimethylaminopyridine (1.22 g, 0.01 mol) and 15.2 g (0.11 mol) of bicyclo[3.2.1]octane-2,4-dione are added at a temperature of from 0 to 5°C. After 3 hours at 22°C, the reaction mixture is extracted with 2 N hydrochloric acid. The methylenechloride phase is separated off, washed with water and then extracted with 10% aqueous sodium bicarbonate solution, dried over sodium sulfate and concentrated. This gives 36.9 g (100% of theory) of 4-oxobicyclo[3.2.1]oct-2-en-2-yl 2-(2-methoxyethoxymethyl)-6-trifluoromethylnicotininate as an oil, which can be used further without purification.

36.9 g (0.1 mol) of 4-oxobicyclo[3.2.1]oct-2-en-2-yl 2-(2-methoxyethoxymethyl)-6-trifluoro-methylnicotinate and 27.9 ml (0.2 mol) of triethylamine are dissolved in 400 ml of acetonitrile. At a temperature of 22°C, 0.92 ml (0.01 mol) of acetone cyanohydrin is added. After 18 hours at 22°C, the reaction mixture is poured into a water/2 N hydrochloric acid mixture and extracted with ethyl acetate. The ethyl acetate phase is washed with water and then with concentrated sodium chloride solution, dried over sodium sulfate and concentrated, and the residue is triturated with hexane. Filtration gives 27.9 g (75.6% of theory) of 4-hydroxyl-3-[2-(2-methoxyethoxymethyl)-6-trifluoromethylpyridine-3-carbonyl]bicyclo[3.2.1]oct-3-en-2-one in the form of white crystals (m.p. 55-56°C).

Example H4: 3-(2-Hydroxy-4-oxobicyclo[3.2.1]oct-2-en-3-carbonyl)-6-trifluoromethylpyridin-2-yl methyl acetate:

5.0 g (1 mmol) of 4-hydroxy-3-(2-methyl-1-oxy-6-trifluoromethylpyridin-3-carbonyl)-bicyclo[3.2.1]oct-3-en-2-one (preparation as described in WO 00/15615) are dissolved in 100 ml of toluene and, in the presence of 6.9 ml (0.073 mol) of acetic anhydride, heated at reflux temperature for 10 hours. The mixture is then partitioned between water and ethyl acetate and the organic phase is dried over sodium sulfate and concentrated under reduced pressure. The residue that remains is chromatographed on silica gel. The viscous oil obtained by eluting with a mixture of toluene, ethyl alcohol, dioxane, triethylamine and water (100:40:20:20:5 parts by volume) is dissolved in ethyl acetate and washed successively with 10% hydrochloric acid and water. The organic solution is dried over Na_2SO_4 and concentrated, giving 2.14 g (38%) of pure 3-(2-hydroxy-4-oxobicyclo[3.2.1]oct-2-ene-3-carbonyl)-6-trifluoromethylpyridin-2-ylmethyl acetate in the form of an oil. ¹H-NMR (250 MHz, CDCl₃): 17.06 (s), 1H; 7.67 (s), 2H; 5.27 (d, J = 12.5 Hz), 1H; 5.20 (d, J = 12.5 Hz), 1H; 3.18, (t, J = 5.0 Hz), 1H; 2.92, (t, J = 5.0 Hz), 1H; 2.29-1.98 (m), 4H; 2.00, (s), 3H; 1.81-1.73 ppm (m), 2H.

<u>Example H5: 4-Hydroxy-3-(2-oxiranylmethoxymethyl-6-trifluoromethylpyridine-3-carbonyl)-bicyclo[3.2.1]oct-3-en-2-one:</u>

5 g (0.013 mol) of 3-(2-hydroxy-4-oxobicyclo[3.2.1]oct-2-ene-3-carbonyl)-6-trifluoromethyl-pyridin-2-ylmethyl acetate are dissolved in 60 ml of methanol/water (3:1 mixture), and 1.4 g (0.046 mol) of lithium hydroxide hydrate are added a little at a time at a temperature of 22°C. After 3 hours at 22°C, the reaction mixture is poured into ethyl acetate and 10% hydrochloric acid, and the organic phase is washed three times with water, dried with sodium sulfate and concentrated. This gives 4.1g of 4-hydroxy-3-(2-hydroxymethyl-6-trifluoromethylpyridine-3-carbonyl)bicyclo[3.2.1]oct-3-en-2-one as an oil which can be reacted further without purification.

1.5 g of 4-hydroxy-3-(2-hydroxymethyl-6-trifluoromethylpyridine-3-carbonyl)-bicyclo[3.2.1]oct-3-en-2-one are dissolved in 15 ml of dimethylformamide and, at room temperature, treated with 0.4 g of sodium hydride (80% suspension in oil, 0.013 mol), a little at a time. After 15 minutes at a temperature of 22°C, 3ml (0.036 mol) of epibromohydrin are added dropwise,

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and the reaction mixture is stirred at this temperature for another 18 hours. Ethyl acetate is then added, and the mixture is acidified to pH 3 using 10% hydrochloric acid and extracted with ethyl acetate. The organic phase is dried over sodium sulfate and the crude product is purified chromatographically (mobile phase: toluene/ethyl alcohol/dioxane/triethylamine/water 100:40:20:20:5 parts by volume). This gives the title compound (triethylamine salt) in the form of a yellowish resin, which is released similarly to example H4. Trituration with hexane gives 600 mg of pure 4-hydroxy-3-(2-oxiranylmethoxy-methyl-6-trifluoromethylpyridin-3-carbonyl)bicyclo[3.2.1]oct-3-en-2-one of melting point 54-56°C.

<u>Example H6: (5-Hydroxy-1,3-dimethyl-1H-pyrazol-4-yl)-[2-(2-methoxyethoxymethyl)-6-trifluoromethylpyridin-3-yl]methanone:</u>

1.0 g (0.004 mol) of 2-(2-methoxyethoxymethyl)-6-trifluoromethylnicotinic acid is dissolved in 10 ml of oxalyl chloride. Three drops of dimethylformamide are added, and the mixture is stirred at room temperature for 1 hour. The mixture is then concentrated using a rotary evaporator, and the residue (2-(2-methoxyethoxymethyl)-6-trifluoromethylnicotinoyl chloride) is taken up in 10 ml of methylene chloride. At a temperature of 0°C, 0.84 ml (0.006 mol) of triethylamine and 0.45 g (0.004 mol) of 2,5-dimethyl-2,4-dihydropyrazol-3-one are added. After 2 hours at a temperature of 22°C, the solvent is removed using a vacuum rotary evaporator, and the residue that remains is dissolved in 10 ml of acetonitrile and, to rearrange the intermediate (2,5-dimethyl-2H-pyrazol-3-yl 2-(2-methoxyethoxymethyl)-6trifluoromethylnicotinate), admixed with 0.1 ml of acetone cyanohydrin and 1.13 ml (0.008 mol) of triethylamine. The reaction solution is stirred at room temperature for four hours and then concentrated. The syrup that remains is chromatographed on silica gel. The viscous oil obtained by eluting with a mixture of toluene, ethyl alcohol, dioxane, triethylamine and water (100:40:20:20:5 parts by volume) is dissolved in ethyl acetate and washed successively with 10% hydrochloric acid and water. The organic solution is dried over Na₂SO₄ and concentrated, giving 0.93 g of (5-hydroxy-1,3-dimethyl-1H-pyrazol-4-yl)-[2-(2methoxyethoxymethyl)-6-trifluoromethylpyridin-3-yl]methanone in the form of a viscous oil. ¹H NMR (300 MHz, CDCl₃, δ in ppm): 7.81, (d, J = 6 Hz), 1H; 7.74, (d, J = 6 Hz), 1H; 4.84, (s), 2H; 2H: 3.71, (s), 3H; 3.59, (t, J = 6 Hz) 2H; 3.38, (dd, J 4.0, 3.0 Hz), 1H; 3.26, (s), 3H; 1.82 ppm, (s), 1H.

Preferred compounds of the formula I and their intermediates are listed in the tables below.

In the table below, the left-hand valency of the radical R_1 is attached to the pyridine ring. If no free valency is indicated in the substituent R_2 , as, for example, in the case of point of attachment is at the "CH" carbon atom.

In the table below, the compounds of the formula I are represented as:

A-Q

where the formula A

$$\begin{array}{c|c} R_4 & O \\ \hline R_3 & N & R_1 \\ \hline (O)p & X_1 & R_2 \end{array} (A)$$

denotes the following radicals:

Radical	R ₁	R ₂	R ₄	R ₃	X ₁	p
A1	CH ₂	CH₃	Н	CF ₃	0	0
A2	CH ₂	CH₂CH₃	Н	CF ₃	0	O
A3	CH₂	(CH ₃) ₂ CH	Н	CF ₃	0	0
A4	CH₂	PhCH ₂	Н	CF₃	0	0
A 5	CH₂ ·	CH₃	Н	CF₃	S	0
A6	CH ₂	CH₃	Н	CF ₃	so	0
A7	CH ₂	CH₃	Н	CF ₃	SO ₂	0
A8	CH₂	CH₃OCH₂	Н	CF ₃	0	0
A9	CH₂	CH₃CH₂OCH₂	Н	CF ₃	0	0
A10	CH₂	CH₃OCH₂CH₂	Н	CF ₃	0	0

Radical	R ₁	R ₂	R ₄	R ₃	X ₁	p
A11	CH ₂	CH ₃ CH ₂ OCH ₂ CH ₂	H	CF ₃	0	$\frac{P}{0}$
A12	CH ₂	CH ₃ OC(CH ₃) ₂ CH ₂	Н	CF₃	0	0
A13	CH₂	CH₃OCH(CH₃)CH₂	Н	CF ₃	0	0
A14	CH ₂	CH ₃ OCH ₂ CH(CH ₃)	Н	CF₃	0	0
A15	CH ₂	CH₃OCH₂C(CH₃)₂	Н	CF₃	0	0
A16	CH ₂	CH₃OCH(CH₃)	н	CF₃	0	. 0
A17	CH₂	CH₃OC(CH₃)₂	Н	CF₃	0	0
A18	CH ₂	HC≡CCH ₂	н	CF₃	0	0
A19	CH₂	H ₂ C=CHCH ₂	Н	CF ₃	0	0
A20	CH ₂	CH₃C≡CCH₂	Н	CF ₃	0	0
A21	CH ₂	СН	Н	CF ₃	0	0
A22	.CH₂	Сн	Н	CF ₃	0	0
A23	CH₂	Ссн	Н	CF ₃	0	0
A24	CH ₂	СН	Н	CF ₃	0	0
A25	CH₂	СН	н	CF ₃	0	0
A26	CH₂	СН	Н	CF ₃	0	0
A27	CH ₂	СН	Н	CF₃	0	0
A28	CH ₂	O_CH	Н	CF₃	0	0
A29	CH ₂	ОСН	Н	CF₃	0	0
A30	CH₂ .	ОСН	н	CF ₃	0	o
A31	CH₂	O CH	Н	CF₃	0	0

Radical	R ₁	R ₂	R ₄	R ₃	X ₁	р
A32	CH₂		Н	CF ₃	0	0
A33	CH₂	OCH3	Н	CF₃	0	0
A34	CH₂	ОН	Н	CF ₃	. 0	0
A35	CH ₂	OCH ₃	Н	CF₃	0	0
A36	CH ₂	OH	Н	CF ₃	0	0
A37	CH₂		Н	CF ₃	0	0
A38	CH₂	CH ₃ CH ₃	Н	CF₃	0	0
A39	CH₂	CH ₃	Н	CF ₃	0	0
A40	CH₂ .	N CH ₃	Н	CF ₃	0	0
A41	CH₂ .	₩ N	Н	CF ₃	Ο	0
A42	CH ₂		Н	CF ₃	0	0
A43	CH₂		Н	CF ₃	0	0

Radical	R ₁	R ₂	R ₄	R ₃	X ₁	p
A44	CH ₂	OCH ₃	Н	CF ₃	0	0
A45	CH ₂	OH N	Н	CF₃	0	0
A46	CH ₂	OCH ₃	н	CF₃	0	0
A47	CH ₂	OH OH	Н	CF₃	0	0
A48	CH ₂	OCH ₃	Н	CF₃	Ο	0
A49	CH ₂	OH OH	Н	CF₃	0	0
A50	CH₂		Н	CF ₃	0	O
A51	CH ₂	⟨ N	Н	CF ₃	0	0
A52	CH ₂	POCH3	Н	CF ₃	0	0
A53	CH₂	OCH ₃	н	CF ₃	0	0
A54	CH₂	CH=CH OCH ₃	Н	CF ₃	0	0

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Radical	R ₁	R ₂	R ₄	R ₃	X ₁	<u>р</u>
A55	CH₂	CH ₂	Н	CF ₃	0	0
A56	CH	OCH ₃				
	CH₂	CH ₂	Н	CF ₃	0	. 0
A 57	CH₂	OCH ₂	Н	CF ₃	Ο	0
A58	CH ₂	CH ₂	Н	CF₃	0	0
A59	CH₂	O CH ₂	Н	CF ₃	0	0
A60	CH₂	CH ₂	н	CF ₃	0	o .
A61	CH ₂	CH ₂	н	CF ₃	0	o
A62	CH₂	CH ₂	н	CF ₃	0	0
A63	CH₂	CH ₂	Н	CF₃	Ο	0
A64	CH₂	CH ₂	н	CF ₃	0	0
A65	CH₂	OCH ₂	Н	CF ₃	. 0	0
A66	CH₂	O CH ₂	Н	CF₃	0	0
A67	CH₂	CH ₂	Н	CF₃	0	0
A68	CH₂	CH ₂ OCH ₃	Н	CF₃	0	0
A69	CH₂	OH OH	Н	CF₃	0	0

Radical	R ₁	R ₂	R ₄	R ₃	X ₁	·p
A70	CH₂	OCH ₃	H	CF₃	0	0
A71	CH₂	CH ₂	Н	CF ₃	. 0	0
A72	CH₂	CH ₂	Н	CF ₃	0	0
A73	CH₂	CH ₃ OCH ₂ CH ₂	Н	CF ₃	0	0
A74	CH ₂	CH ₃ OCH ₂ CH ₂	Н	CF₃	0	0
A75	CH ₂	OCH ₂ CH ₂	Н	CF₃	0	0
A76	CH ₂	CH ₂	н	CF ₃	0	0
A77	CH ₂	CH ₂	Н	CF ₃	0	0
A78	CH ₂	CH ₂	н	CF ₃	0	0
A79	· CH₂	OCH ₃ CH ₂	H	CF ₃	0	0
A80	CH₂ ·	OH CH ₂	Н	CF₃	0	0
A81	CH₂	OCH ₃ CH ₂	Н	CF₃	0	0

Radical	R ₁	R ₂	R ₄	R ₃	X ₁	р
A82	CH₂	OH CH₂	Н	CF₃	0	0
A83	ĊH₂	OCH ₃ CH ₂	Н	CF₃	0	0
A84	CH₂	OH CH ₂	Н	CF ₃	0	0
· A85	CH₂	CH ₂	Н	CF₃	0	0
A86	CH₂	CH ₂	Н	CF₃	0	O
A87	CH₂	F OCH ₃	Н	CF₃	0	0
A88	CH₂	OCH ₂ CH ₂	Н	CF ₃	0	0
A89	CH₂	OCH ₂ CH ₂	Н	CF₃	0	Ò
A90	CH₂	OCH ₃	Н	CF₃	0	0
A91	CH₂CH₂	CH₃	Н	CF ₃	0	0
A92	CH ₂ CH ₂	CH₃CH₂	Н	CF₃	Ο.	0
A93	CH₂CH₂	(CH ₃) ₂ CH	Н	CF ₃	0	0
A94	CH₂CH₂ [′]	PhCH₂	Н	CF ₃	0	0
A95	CH₂CH₂	CH ₃	Н	CF ₃	s	0
A96	CH₂CH₂	CH ₃	Н	CF₃	so	0
A97	CH₂CH₂	CH ₃	Н	CF ₃	SO ₂	0
A98	CH ₂ CH ₂	(CH ₃) ₂ CHCH ₂	Н	CF ₃	0	

Radical	R ₁	R ₂	R ₄	R ₃	Х ₁	р
A99	CH ₂ CH ₂	CH₃OCH₂	Н	CF ₃	0	0
A100	CH₂CH₂	CH₃CH₂OCH₂	Н	CF ₃	О	0
A101	CH₂CH₂	CH₃OCH₂CH₂	Н	CF ₃	0	0
A102	CH ₂ CH ₂	CH₃CH₂OCH₂CH₂	Н	CF ₃	0	0
A103	CH ₂ CH ₂	CH ₃ OC(CH ₃) ₂ CH ₂	Н	CF₃	0	0
A104	CH ₂ CH ₂	CH ₃ OCH(CH ₃)CH ₂	Н	CF ₃	0	0
A105	CH₂CH₂	CH₃OCH₂CH(CH₃)	Н	CF ₃	0	0
A106	CH₂CH₂	CH ₃ OCH ₂ C(CH ₃) ₂	Н	CF ₃	0	0
A107	CH ₂ CH ₂	CH₃OCH(CH₃)	Н	CF₃	0	0
A108	CH₂CH₂	CH₃OC(CH₃) ₂	Н	CF ₃	0	0
A109	CH₂CH₂	HC≡CCH ₂	Н	CF ₃	Ο	0
A110	CH ₂ CH ₂	H ₂ C=CHCH ₂	Н	CF ₃	0	0
A111	CH₂CH₂	CH₃C≡CCH₂	Н	CF ₃	O	0
A112	CH₂CH₂	СН	Н	CF ₃	0	0
A113	CH₂CH₂	Сн	Н	CF ₃	0	0
A114	CH₂CH₂	Ссн	Н	CF ₃	0	0
A115	CH ₂ CH ₂	СН	Н	CF ₃	Ο	.0
A116	CH₂CH₂	СН	Н	CF₃	0	0
A117	CH₂CH₂	СН	Н	CF₃	0	0
A118	CH ₂ CH ₂	СН	Н	CF₃	Ο	0
A119	CH₂CH₂	O_CH	Н	CF ₃	0	0
A120	CH ₂ CH ₂	O_CH	Н	CF₃	Ο	0
A121	CH₂CH₂	ОСН	Н	CF ₃	0	0

Radical	R ₁	R ₂	R ₄	R ₃	X ₁	р
A122	CH₂CH₂	O CH	Н	CF ₃	O	0
A123	CH₂CH₂		Н	CF ₃	O .	0
A124	CH₂CH₂	OCH ₃	Н	CF₃	0	0
A125	CH₂CH₂	ОН	Н	CF ₃	Ο	0
A126	CH₂CH₂		Н	CF₃	0	0
A127	CH₂CH₂	ÓCH₃	н	CF₃	0	Ŏ
A128	CH ₂ CH ₂	óн	н	CF ₃	0	0
A129	CH₂CH₂	CH ₃ CH ₃	н	CF ₃	0	0
A130	CH ₂ CH ₂	CH ³	н	CF₃	0	0
A131	CH₂CH₂	cH₃ NNN CH₃	H	CF₃	O	0
A132	CH₂CH₂ ·	CH₃	Н	CF ₃	0	0
A133	CH₂CH₂		Н	CF₃	Ο	0

Radical	R ₁	R ₂	R ₄	R ₃	X ₁	р
A134	CH₂CH₂	N	Н	CF ₃	0	0
A135	CH₂CH₂	OCH ₃	Н	CF ₃	0	0
A136	CH₂CH₂	OH N	Н	CF₃	0	0
A137	CH₂CH₂	OCH ₃	Н	CF₃	0	0
A138	CH₂CH₂	OH	Н	CF₃	0	0
A139	CH₂CH₂	OCH ₃	Н	CF₃	0	0
A140	CH₂CH₂	OH	Н	CF₃	0	0
A141	CH ₂ CH ₂		Н	CF ₃	O	0
A142	CH₂CH₂	N	н	CF ₃	0	0
A143	CH₂CH₂	POCH3	Н	CF ₃	0	
A144	CH₂CH₂	OCH ₃	Н	CF ₃	0	0

Radical	R ₁	R ₂	R ₄	R ₃	X ₁	р
A145	CH₂CH₂	CH ₂ CH=CH OCH ₃	Н	CF₃	0	0
A146	CH₂CH₂	CH ₂	Н	CF₃	0	0
A147	CH ₂ CH ₂	CH ₂	Н	CF ₃	Ο	0
A148	CH ₂ CH ₂	O CH ₂	Н	CF₃	0	0
A149	CH₂CH₂	CH ₂	Н	CF ₃	0	0
A150	CH₂CH₂	CH₂	Н	CF ₃	0	0
A151	CH₂CH₂	CH ₂	Н	CF ₃	0	0
A152	CH₂CH₂	CH ₂	Н	CF₃	0	0
A153	CH ₂ CH ₂	CH ₂	Н	CF ₃	0	0
A154	CH₂CH₂	O CH ₂	Н	CF ₃	O	0
A155	CH₂CH₂	CH ₂	Н	CF₃	Ο	0
A156	CH ₂ CH ₂	OCH ₂	Н	CF₃	0	0
A157	CH₂CH₂ .	O CH ₂	Н	CF ₃	0	0
A158	CH₂CH₂	CH ₂	Н	CF₃	0	0

Radical	R ₁	R ₂	R ₄	R ₃	X ₁	p
A159	CH₂CH₂	CH ₂ OCH ₃	H .	CF ₃	0	0
A160	CH₂CH₂	CH₂ OH	Н	CF₃	0	0
A161	CH₂CH₂	CH ₂ OCH ₃	Н	CF₃	0	0
A162	CH ₂ CH ₂	CH ₂	Н	CF₃	0	0
A163	CH₂CH₂	CH ₂	н	CF₃	0	0
A164	CH₂CH₂	CH ₃ OCH ₂ CH ₂	Н	CF₃	0	0
A165	CH ₂ CH ₂	CH ₃ N OCH ₂ CH ₂	н	CF₃	0	0
A166	CH ₂ CH ₂	NNOCH ₂ CH ₂	Н	CF₃	0	Ö
A167	CH ₂ CH ₂	CH ₂	Н	CF₃	O	0
A168	CH₂CH₂	CH ₂	Н	CF₃	Ó	0
A169	CH₂CH₂ ·	CH ₂	Н	CF₃	0	0
A170	CH₂CH₂	OCH ₃ CH ₂	Н	CF ₃	0	0

Radical	R ₁	R ₂	R ₄	R ₃	X ₁	р
A171	CH₂CH₂	OH CH ₂	Н	CF ₃	0	0
A172	CH₂CH₂	OCH ₃	Н	CF₃	0	0
A173	CH₂CH₂	OH CH₂	Н	CF₃	0	0
A174	CH₂CH₂	OCH ₃ CH ₂	Н	CF ₃	0	0
A175	CH ₂ CH ₂	OH CH ₂	Н	CF₃	0	0
A176	CH₂CH₂	CH ₂	Н	CF₃	Ο	0
A177	· CH₂CH₂	CH ₂	Н	CF ₃	0	0
A178	CH₂CH₂	F CH ₂ OCH ₃	Н	CF₃	0	0
A179	CH₂CH₂	OCH ₂ CH ₂	Н	CF ₃	0	0
A180	CH₂CH₂	OCH ₂ CH ₂	Н	CF ₃	0	0
A181	CH₂CH₂	OCH ₂	Н	CF₃	0	0
A182	CH(OCH₃)CH₂	CH ₃	Н	CF ₃	0	0
A183	CH(OCH₃)CH₂	CH₃CH₂	Н	CF ₃	0	0
A184	CH(OCH ₃)CH ₂	(CH₃)₂CH	Н	CF ₃	0	0

Radical	R ₁	R ₂	R ₄	R ₃	Х1	p
A185	CH(OCH₃)CH₂	PhCH₂	Н	CF ₃	0	0
A186	CH(OCH₃)CH₂	CH₃	Н	CF ₃	S	0
A187	CH(OCH ₃)CH ₂	CH₃	Н	CF ₃	SO	0
A188	CH(OCH₃)CH₂	CH₃	Н	CF ₃	SO ₂	0
A189	CH(OCH₃)CH₂	CH₃CH₂CH₂	Н	CF ₃	Ο	0
A190	CH(OCH₃)CH₂	CH₃OCH₂	Н	CF ₃	0	0
A191	CH(OCH ₃)CH ₂	CH₃CH₂OCH₂	Н	CF ₃	0	0
A192	CH(OCH ₃)CH ₂	CH₃OCH₂CH₂	Н	CF₃	0	0
A193	CH(OCH₃)CH₂	CH₃CH₂OCH₂CH₂	Н	CF ₃	0	0
A194	CH(OCH₃)CH₂	CH₃OC(CH₃)₂CH₂	·H	CF ₃	0	0
A195	CH(OCH₃)CH₂	CH₃OCH(CH₃)CH₂	Н	CF ₃	0	0
A196	CH(OCH₃)CH₂	CH ₃ OCH ₂ CH(CH ₃)	Н	CF ₃	0	0
A197	CH(OCH ₃)CH ₂	CH ₃ OCH ₂ C(CH ₃) ₂	Н	CF ₃	0	0
A198	CH(OCH₃)CH₂	CH₃OCH(CH₃)	Н	CF ₃	0	0
A199	CH(OCH₃)CH₂	CH ₃ OC(CH ₃) ₂	Н	CF ₃	0	0
A200	CH(OCH₃)CH₂	HC≡CCH ₂	Н	CF ₃	0	0
A201	CH(OCH₃)CH₂	H₂C=CHCH₂	Н	CF ₃	0	0
A202	CH(OCH₃)CH₂	CH₃C≡CCH₂	Н	CF ₃	0	0
A203	CH(OCH₃)CH₂	Сн	Н	CF₃	0	0
A204	CH(OCH₃)CH₂	Осн	Н	CF ₃	0	0
A205	CH(OCH₃)CH₂	Ссн	Н	CF ₃	0	0
A206	CH(OCH ₃)CH ₂	° CH	Н	CF ₃	Ο	0
A207	CH(OCH₃)CH₂	Сн	Н	CF ₃	0	0
A208	CH(OCH₃)CH₂	СН	Н	CF ₃	0	0
A209	CH(OCH₃)CH₂	СН	Н	CF ₃	0	0
A210	CH(OCH₃)CH₂	ОСН	H.	CF₃	0	0

Radical	R ₁	R ₂	R ₄	R ₃	X ₁	р
A211	CH(OCH₃)CH₂	O_CH	Н	CF ₃	0	0
A212	CH(OCH₃)CH₂	ОСН	н	CF ₃	0	0
A213	CH(OCH₃)CH₂	O_CH	Н	CF ₃	0	0
A214	CH(OCH₃)CH₂		H	CF₃	O´	0
A215	CH(OCH₃)CH₂	OCH ₃	Н	CF ₃	0	0
A216	CH(OCH₃)CH₂	ОН	Н	CF₃	0	0
A217	CH(OCH₃)CH₂		Н	CF ₃	Ο.	0
A218	CH(OCH₃)CH₂	о́сн₃	Н	CF ₃	0	0
A219	CH(OCH₃)CH₂	он	Н	CF ₃	0	0
A220	CH(OCH₃)CH₂	CH ₃ CH ₃	н	CF₃	Ο	, 0
A221	CH(OCH₃)CH₂	CH ₃	Н	CF₃	0	0
A222	CH(OCH₃)CH₂	NNN CH3	·H	CF ₃	0	0

Radical	R ₁	R ₂	R ₄	R ₃	Х ₁	p
A223	CH(OCH ₃)CH ₂	N N	Н	CF ₃	0	0
A224	CH(OCH₃)CH₂		Н	CF₃	0	0
A225	CH(OCH₃)CH₂	N N	Н	CF₃	0	0
A226	CH(OCH₃)CH₂	OCH ₃	Н	CF₃	0	0
A227	CH(OCH₃)CH₂	OH N	Н	CF₃	0	0
A228	CH(OCH₃)CH₂	OCH ₃	Н	CF₃	0	0
A229	CH(OCH₃)CH₂	OH	Н	CF ₃	0	0
A230	CH(OCH₃)CH₂	OCH ₃	н	CF ₃	0	0
A231	CH(OCH₃)CH₂	OH N	Н	CF ₃	0	0
A232	CH(OCH₃)CH₂		Н	CF ₃	0	0
A233	CH(OCH₃)CH₂		Н	CF ₃	0	0
A234	CH(OCH₃)CH₂	F OCH3	Н	CF ₃	Ο	0

Radical	R ₁	R ₂	R ₄	R ₃	X ₁	p
A235	CH(OCH₃)CH₂	NOCH ₃	Н	CF ₃	0	0
A236	CH(OCH₃)CH₂	CH=CH OCH ₃	н	CF₃	0	0
A237	CH(OCH₃)CH₂	OCH ₃	Н	CF₃	0	0
A238	CH(OCH₃)CH₂	CH ₂	Н	CF ₃	0	0
A239	CH(OCH₃)CH₂	$^{\rm O}$ $^{\rm CH_2}$	Н	CF ₃	O	0
A240	CH(OCH₃)CH₂	CH ₂	Н	CF ₃	0	0
A241	CH(OCH₃)CH₂	O CH ₂	Н	CF ₃	0	0
A242	CH(OCH₃)CH₂	CH ₂	Н	CF ₃	0	0
A243	CH(OCH₃)CH₂	CH ₂	Н	CF ₃	0	0
A244	CH(OCH₃)CH₂	CH ₂	Н	CF₃	0	0
A245	CH(OCH₃)CH₂	CH ₂	Н	CF ₃	0	0
A246	CH(OCH₃)CH₂	CH ₂	Н	CF₃	0	0
A247	CH(OCH₃)GH₂	OCH ₂	Н	CF ₃	0	0
A248	CH(OCH₃)CH₂	O CH ₂	Н	CF₃	0	0

Radical	R ₁	R ₂	R ₄	R ₃	X ₁	p
A249	CH(OCH₃)CH₂	CH ₂	Н	CF₃	0	0
A250	CH(OCH₃)CH₂	CH ₂ OCH ₃	Н	CF₃	0	0
A251	CH(OCH₃)CH₂	CH₂ OH	Н	CF ₃	0	0
A252	CH(OCH₃)CH₂	CH ₂ OCH ₃	Н	CF ₃	0	0
A253	CH(OCH₃)CH₂	OH CH ₂	Н	CF ₃	0	0
A254	CH(OCH₃)CH₂	CH₂ S	н	CF ₃	0	0
A255	CH(OCH₃)CH₂	CH ₃ OCH ₂ CH ₂	Н	CF ₃	0	0
A256	CH(OCH₃)CH₂	CH ₃ N N OCH₂CH₂ CH₃	Н	CF₃	Ο	0
A257	CH(OCH₃)CH₂	OCH ₂ CH ₂	Н	CF₃	0	0
A258	CH(OCH₃)CH₂	CH ₂	Н	CF₃	0	0
A259	CH(OCH₃)CH₂	CH ₂	Н	CF₃	0	0
A260	CH(OCH₃)CH₂	CH ₂	Н	CF₃	0	0

Radical	R ₁	R ₂	R ₄	R ₃	X ₁	p
A261	CH(OCH₃)CH₂	OCH ₃ CH ₂	Н	CF ₃	0	0
A262	CH(OCH₃)CH₂	OH CH ₂	Н	CF₃	0	0
A263	CH(OCH₃)CH₂	OCH ₃ CH ₂	Н	CF ₃	0	0
A264	CH(OCH₃)CH₂ ·	OH CH ₂	Н	CF ₃	0	0
A265	CH(OCH₃)CH₂	OCH ₃ CH ₂	Н	CF₃	0	0
A266	CH(OCH₃)CH₂	OH CH ₂	Н	CF₃	Ο	0
A267	CH(OCH₃)CH₂	CH ₂	Н	CF₃	0	O
A268	CH(OCH₃)CH₂	CH ₂	Н	CF ₃	0	0
A269	CH(OCH₃)CH₂	F OCH ₂	H	CF₃	0	0
A270	CH(OCH₃)CH₂	OCH ₂ CH ₂	Н	CF₃	0	0
A271	CH(OCH₃)CH₂	OCH ₂ CH ₂	Н	CF₃	0	0
A272	CH(OCH₃)CH₂	OCH ₂	Н.	CF ₃	0	0

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Radical	R ₁	R ₂	R ₄	R ₃	X ₁	р
A273	CH ₂ CH(OCH ₃)CH ₂	CH₃	Н	CF ₃	0	0
A274	CH₂CH(OCH₃)CH₂	CH₃CH₂	Н	CF ₃	0	0
A275	CH₂CH(OCH₃)CH₂	(CH₃)₂CH	Н	CF ₃	0	0
A276	CH₂CH(OCH₃)CH₂	PhCH₂	Н	CF ₃	0	0
A277	CH₂CH(OCH₃)CH₂	CH₃	Н	CF ₃	S	0
A278	CH₂CH(OCH₃)CH₂	CH₃	Н	CF ₃	so	0
A279	CH₂CH(OCH₃)CH₂	CH₃	Н	CF3	SO ₂	0
A280	CH ₂ CH(OCH ₃)CH ₂	CH ₃ CH ₂ CH ₂	Н	CF ₃	0	0
A281	CH₂CH(OCH₃)CH₂	CH₃OCH ₂	Н	CF ₃	0	0
A282	CH₂CH(OCH₃)CH₂	CH3CH2OCH2	Н	CF ₃	0	0
A283	CH₂CH(OCH₃)CH₂	CH₃OCH₂CH₂	Н	CF ₃	0	0
A284	CH₂CH(OCH₃)CH₂	CH₃CH₂OCH₂CH₂	Н	CF ₃	0	0.
A285	CH₂CH(OCH₃)CH₂	CH ₃ OC(CH ₃) ₂ CH ₂	Н	CF ₃	0	0
A286	CH₂CH(OCH₃)CH₂	CH ₃ OCH(CH ₃)CH ₂	Н	CF ₃	0	0
A287	CH₂CH(OCH₃)CH₂	CH₃OCH₂CH(CH₃)	Н	CF ₃	0	0
A288	CH₂CH(OCH₃)CH₂	CH₃OCH₂C(CH₃)₂	Н	CF ₃	0	0
A289	CH₂CH(OCH₃)CH₂	CH₃OCH(CH₃)	Н	CF ₃	0	0
A290	CH₂CH(OCH₃)CH₂	CH ₃ OC(CH ₃) ₂	Н	CF ₃	0	0
A291	CH₂CH(OCH₃)CH₂	HC≡CCH ₂	Н	CF ₃	0	0
A292	CH ₂ CH(OCH ₃)CH ₂	H ₂ C=CHCH ₂	Н	CF ₃	0	0
A293	CH ₂ CH(OCH ₃)CH ₂	CH ₃ C≡CCH ₂	Н	CF ₃	0	0
A294	CH₂CH(OCH₃)CH₂	Сн	Н	CF ₃	Ο.	0
A295	CH₂CH(OCH₃)CH₂	Осн	Н	CF ₃	0	0
A296	CH₂CH(OCH₃)CH₂	Ссн	н	CF ₃	0	0
A297	CH₂CH(OCH₃)CH₂	o√ch	Н	CF₃	0	0
A298	CH₂CH(OCH₃)CH₂	СН	Н	CF ₃	0	0
A299	CH₂CH(OCH₃)CH₂	СН	Н	CF₃	0	0

Radical		R ₂	R ₄	R ₃	X ₁	р
A300	CH₂CH(OCH₃)CH₂	СН	Н	CF ₃	0	0
A301	CH₂CH(OCH₃)CH₂	O_CH	Н	CF ₃	0	0
A302	CH₂CH(OCH₃)CH₂	O_CH	Ĥ	CF ₃	0	0
A303	CH ₂ CH(OCH ₃)CH ₂	СН	Н	CF ₃	0	0
A304	CH ₂ CH(OCH ₃)CH ₂	O_CH	Н	CF₃	0	0
A305	CH₂CH(OCH₃)CH₂		Н	CF₃	0	0
A306	CH ₂ CH(OCH ₃)CH ₂	OCH ₃	Н	CF ₃	0	0
A307	CH₂CH(OCH₃)CH₂	ОН	Н	CF₃	O .	0
A308	CH₂CH(OCH₃)CH₂	OCH ₃	Н	CF ₃	0	p ·
A309	CH₂CH(OCH₃)CH₂	OH	н	CF ₃	0	
A310	CH₂CH(OCH₃)CH₂		Н	CF ₃	0	0
A311	CH₂CH(OCH₃)CH₂	CH ₃ CH ₃	Н	CF ₃	Ο	0
A312	CH₂CH(OCH₃)CH₂	CH ₃	Н	CF ₃	O	0

Radical	R ₁	R ₂	R ₄	R ₃	X ₁	р
A313	CH₂CH(OCH₃)CH₂	N N CH ₃	Н	CF ₃	0	0
A314	CH₂CH(OCH₃)CH₂		Н	CF ₃	. О	0
A315	CH₂CH(OCH₃)CH₂		Н	CF₃	0	0
A316	CH ₂ CH(OCH ₃)CH ₂		Н	CF ₃	0	0
A317	CH₂CH(OCH₃)CH₂	OCH ₃	Н	CF ₃	0	0
A318	CH₂CH(OCH₃)CH₂	OH N	Н	CF₃	0	0
A319	CH₂CH(OCH₃)CH₂	NOCH3	H	CF₃	Ο	0
A320	CH₂CH(OCH₃)CH₂	OH OH	Н	CF ₃	0	0
A321	CH₂CH(OCH₃)CH₂	OCH ₃	н	CF ₃	0	0
A322	CH₂CH(OCH₃)CH₂	OH N	H	CF ₃	O	0
A323	CH₂CH(OCH₃)CH₂	ON.	н	CF ₃	0	0
A324	CH₂CH(OCH₃)CH₂	(N	н	CF ₃	0	0

Radical		R ₂	R ₄	R ₃	X ₁	р
A325	CH₂CH(OCH₃)CH₂	POCH ₃	Н	CF₃	0	0
A326	CH₂CH(OCH₃)CH₂	OCH ₃	Н	CF₃	0	0
A327	CH₂CH(OCH₃)CH₂	CH ₂ CH=CH OCH ₃	Н	CF₃	0	0
A328	CH₂CH(OCH₃)CH₂	CH ₂	Н	CF ₃		0
A329	CH ₂ CH(OCH ₃)CH ₂	CH ₂	Н	CF ₃	0	0
A330	CH ₂ CH(OCH ₃)CH ₂	O CH₂	Н	CF ₃	0	0
A331	CH ₂ CH(OCH ₃)CH ₂	CH ₂	Н	CF₃	0	0
A332	CH ₂ CH(OCH ₃)CH ₂	O CH ₂	Н	CF ₃	0	0
A333	CH ₂ CH(OCH ₃)CH ₂	CH ₂	Н	CF ₃	0	0
A334	CH₂CH(OCH₃)CH₂	CH ₂	Н	CF ₃		0
A335	CH₂CH(OCH₃)CH₂	CH ₂	Н	CF₃	0	0
A336	CH₂CH(OCH₃)CH₂	O CH ₂	Н	CF ₃	0	0
A337	CH₂CH(OCH₃)CH₂	CH ₂	H	CF ₃	0	0
A338	CH₂CH(OCH₃)CH₂	OCH ₂	Н	CF₃	0	0

Radical	R ₁	R ₂	R ₄	R ₃	X ₁	р
A339	CH₂CH(OCH₃)CH₂	O CH ₂	Н	CF ₃	0	0
A340	CH ₂ CH(OCH ₃)CH ₂	CH ₂	Н	CF ₃	Ο	Ó
A341	CH₂CH(OCH₃)CH₂	OCH ₃	Н	CF₃	0	0
A342	CH₂CH(OCH₃)CH₂	CH₂ OH	Н	CF₃	0	0
A343	CH₂CH(OCH₃)CH₂	CH ₂	Н	CF ₃	0	0
A344	CH ₂ CH(OCH ₃)CH ₂	CH ₂	н	CF ₃	0	0
A345	CH ₂ CH(OCH ₃)CH ₂	CH₂ S	н	CF ₃	0	0
A346	CH₂CH(OCH₃)CH₂	CH ₃ OCH ₂ CH ₂	Н	CF₃	O	0
A347	CH₂CH(OCH₃)CH₂	CH ₃ N N OCH ₂ CH ₂	н	CF ₃	0	0
A348	CH₂CH(OCH₃)CH₂	CH ₃ OCH ₂ CH ₂	н	CF ₃	0	0
A349	CH₂CH(OCH₃)CH₂	CH ₂	Н	CF ₃	0	0
A350	CH₂CH(OCH₃)CH₂	CH ₂	н	CF₃	Ο	0

Radical	·	R ₂	R ₄	R ₃	X ₁	р
A351	CH₂CH(OCH₃)CH₂	CH ₂	Н	CF₃	0	0
A352	CH₂CH(OCH₃)CH₂	OCH ₃ CH ₂	Н	CF ₃	O	0
A353	CH ₂ CH(OCH ₃)CH ₂	OH CH ₂	Н	CF₃	. О	. 0
A354	CH ₂ CH(OCH ₃)CH ₂	OCH ₃ CH ₂	Н	CF ₃	0	0
A355	CH₂CH(OCH₃)CH₂	OH CH ₂	Н	CF ₃	0	0
A356	CH₂CH(OCH₃)CH₂	OCH ₃ CH ₂	Н	CF ₃	Ο	0
A357	CH₂CH(OCH₃)CH₂	OH CH ₂	Н	CF ₃	Ο	0
A358	CH₂CH(OCH₃)CH₂	CH ₂	Н	CF ₃	0	0
A359	CH₂CH(OCH₃)CH₂	CH ₂	Н	CF ₃	0	0
A360	CH₂CH(OCH₃)CH₂	F CH₂ OCH₃	Ĥ	CF₃	0	0
A361	CH₂CH(OCH₃)CH₂	OCH ₂ CH ₂	Н	CF₃	Ο	0
A362	CH₂CH(OCH₃)CH₂	OCH ₂ CH ₂	Н	CF₃	Ο	0

Radical	R ₁	R ₂	. R ₄	R ₃	X ₁	р
A363	CH₂CH(OCH₃)CH₂	O_CH ₂	Н	CF ₃	0	0
		OCH ₃				
A364	CH=CHCH₂	CH₃	Н	CF ₃	0	0
A365	CH=CHCH ₂	CH ₃ CH ₂	Н	CF ₃	0	0
A366	CH=CHCH ₂	(CH ₃) ₂ CH	Н	CF ₃	0	0
A367	CH=CHCH ₂	PhCH₂	Н	CF ₃	0	0
A368	CH=CHCH₂	CH₃	Н	CF ₃	\$	0
A369	CH=CHCH₂	CH₃	Н	CF ₃	so	0
A370	CH=CHCH₂	CH₃	Н	CF ₃	SO ₂	0
A371	CH=CHCH₂	CH₃CH₂CH₂	н	CF ₃	0	0
A372	CH=CHCH₂	CH₃OCH₂	Н	CF ₃	0	0
A373	CH=CHCH₂	CH₃CH₂OCH₂	Н	CF ₃	0	0
A374	CH=CHCH₂	CH₃OCH₂CH₂	Н	CF₃	0	0
A375	CH=CHCH₂	CH3CH2OCH2CH2	Н	CF ₃	0	0
A376	CH=CHCH₂	CH ₃ OC(CH ₃) ₂ CH ₂	н	CF ₃	0	0
A377	CH=CHCH₂	CH₃OCH(CH₃)CH₂	Н	CF₃	0	0
A378	CH=CHCH ₂	CH₃OCH₂CH(CH₃)	Н	CF ₃	0	0
A379	CH=CHCH ₂	CH ₃ OCH ₂ C(CH ₃) ₂	H	CF₃	0	O
A380	CH=CHCH ₂	CH₃OCH(CH₃)	Н	CF ₃	0	0
A381	CH=CHCH ₂	CH ₃ OC(CH ₃) ₂	Н	CF ₃	0	0
A382	CH=CHCH₂	HC≡CCH ₂	Н	CF ₃	0	0
A383	CH=CHCH₂	H ₂ C=CHCH ₂	Н	CF₃	0	0
A384	CH=CHCH₂	CH₃C≡CCH₂	Н	CF₃	0	0
A385	CH=CHCH₂	Сн	Н	CF ₃	0	0
A386	CH=CHCH₂	Осн	Н	CF ₃	0	0
A387	CH=CHCH₂	CH	Н	CF ₃	0	0
A388	CH=CHCH₂	√ CH	Н	CF ₃	0	0
A389	CH=CHCH ₂	Ссн	Н	CF ₃	0	0

Radical	R ₁	R ₂	R ₄	R ₃	X ₁	р
A390	CH=CHCH₂	СН	Н	CF₃	0	0
A391	CH=CHCH₂	СН	Н	CF₃	0	0
A392	CH=CHCH₂	O_CH	н	CF ₃	0	0
A393	CH=CHCH₂	O_CH	Н	CF ₃	0	0
A394	CH=CHCH₂	ОСН	Н	CF₃	0	0
A395	CH=CHCH₂	ОСН	Н	CF ₃	. 0	0
A396	CH=CHCH₂		Н	CF ₃	0	0
A397	CH=CHCH₂	OCH ₃	Н	CF ₃	0	0
A398	CH=CHCH ₂	ОН	Н	CF₃	0	0
A399	CH=CHCH ₂	OCH ₃	Н	CF₃	0	0
A400	CH=CHCH₂	OH	Н	CF₃	0	0
A401	CH=CHCH₂		н	CF₃	0	0
A402	CH=CHCH₂	CH ₃ CH ₃	Н	CF₃	Ο	0

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Radical	R ₁	R ₂	R ₄	R ₃	X ₁	р
A403	CH=CHCH₂	CH ₃	Н	CF ₃	0	0
A404	CH=CHCH₂	NNN CH ₃	Н	CF₃	0	0
A405	CH=CHCH ₂		Н	CF ₃	0	0
A406	CH=CHCH₂		Н	CF ₃	Ó	0
A407	CH=CHCH ₂		Н	CF ₃	Ο,	0
A408	CH=CHCH₂	OCH ₃	Н	CF₃	Ο	0
A409	CH=CHCH ₂	OH N	Н	CF₃	Ο	0
A410	CH=CHCH₂	N OCH3	н	CF ₃	0	Ò
A411	CH=CHCH₂	OH	Н	CF ₃	0	0
A412	CH=CHCH₂	OCH ₃	Н	CF ₃	O	0
A413	CH=CHCH₂	OH	Н	CF₃	0	0
A414	CH=CHCH₂		Н	CF₃	0	0

Radical	R ₁	R ₂	R ₄	R ₃	X ₁	р
A415	CH=CHCH₂	N N	Н	CF₃	0	0
A416	CH=CHCH₂	POCH3	Н	CF₃	0	0
A417	CH=CHCH₂	OCH ₃	Н	CF₃	0	0 :
A418	CH=CHCH₂	CH=CH OCH ₃	Н	CF₃	0	0
A419	CH=CHCH ₂	CH ₂	Н	CF₃	0	0
A420	CH=CHCH₂	СH ₂	Н	CF ₃	0	0
A421	CH=CHCH₂	. O	н	CF ₃	0	0
A422	CH=CHCH₂	CH ₂	Н	CF ₃	0	0
A423	CH=CHCH ₂	O CH ₂	Н	CF ₃	0	0
A424	CH=CHCH₂	CH ₂	Н	CF ₃	0	0
A425	CH=CHCH₂	CH ₂	Н	CF ₃	0	0
A426	CH=CHCH₂	CH ₂	Н	CF₃	0	0
A427	CH=CHCH₂	O CH ₂	н	CF₃	0	0
A428	CH=CHCH₂	CH ₂	н	CF₃	0	0

Radical	R ₁	R ₂	R ₄	R ₃	X ₁	p
A429	CH=CHCH₂	OCH ₂	Н	CF ₃	0	0
A430	CH=CHCH₂	O CH ₂	н	CF ₃	0	0
A431	CH=CHCH₂	CH ₂	Н	CF₃	0	0
A432	CH=CHCH₂	CH ₂ OCH ₃	н	CF ₃	0	0
A433	CH=CHCH ₂	CH ₂	Н	CF₃	0	0
A434	CH=CHCH ₂	CH ₂ OCH ₃	Н	CF ₃	0	0
A435	CH=CHCH₂	CH ₂	Н	CF₃	O	0
A436	CH=CHCH ₂	CH ₂	Н	CF₃	0	0
A437	CH=CHCH₂	CH ₃ OCH ₂ CH ₂	Н	CF ₃	0	0
A438	CH=CHCH₂	CH ₃ N N OCH ₂ CH ₂	н	CF ₃	0	0
A439	CH=CHCH₂	OCH ₂ CH ₂	н	CF ₃	0	0
A440	CH=CHCH₂	CH ₂	Н	CF ₃	0	0

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Radical	R ₁	R ₂	R ₄	R ₃	X ₁	р
A441	CH=CHCH₂	CH ₂	Н	CF₃	0	0
A442	CH=CHCH ₂	CH₂ N	Н	CF ₃	0	0
A443	CH=CHCH ₂	OCH ₃ CH ₂	H,	CF ₃	0	0
A444	CH=CHCH₂	OH CH ₂	Н	CF₃	0	0
A445	CH=CHCH₂	OCH ₃ CH ₂	Н	CF ₃	0	0
A446	CH=CHCH₂	OH CH ₂	H	CF ₃	0	0
A447	CH=CHCH₂	OCH ₃ CH ₂	Н	CF ₃	Ο	0
A448	CH=CHCH₂	OH CH₂	Н	CF₃	0	0
A449	CH=CHCH₂	CH ₂	Н	CF ₃	0	O
A450	CH=CHCH₂	CH ₂	Н	CF ₃	0	0
A451	CH=CHCH ₂	CH ₂ OCH ₃	н	CF ₃	0	0
A452	CH=CHCH₂	OCH ₂ CH ₂	Н	CF ₃	0	0

Radical	R ₁	R ₂	R ₄	R ₃	X ₁	р
A453	CH=CHCH₂	OCH ₂ CH ₂	Н	CF ₃	0	0
A454	CH=CHCH₂	OCH ₂	Н	CF₃	0	0
A455	C≡CCH ₂	CH₃	Н	CF ₃	0	0
A456	C≡CCH ₂	CH₃CH₂	Н	CF₃	0	0
A457	C≡CCH ₂	(CH₃)₂CH	Н	CF ₃	0	0
A458	C≡CCH₂	PhCH₂	Н	CF ₃	0	0
A459	C≡CCH ₂	CH₃	Н	CF ₃	S	0
A460	C≡CCH ₂	CH₃	Н	CF ₃	SO	0
A461	C≡CCH ₂	CH₃	Н	CF ₃	SO ₂	0
A462	C≡CCH ₂	CH₃CH₂CH₂	Н	CF ₃	0	0
A463	C≡CCH ₂	CH₃OCH ₂ _	Н	CF ₃	0	0
A464	C≡CCH ₂	CH₃CH₂OCH₂	Н	CF ₃	0	0
A465	C≡CCH ₂	CH₃OCH₂CH₂	Н	CF ₃	0	0
A466	C≡CCH ₂	CH ₃ CH ₂ OCH ₂ CH ₂	Н	CF ₃	Ο	0
A467	C≡CCH ₂	CH ₃ OC(CH ₃) ₂ CH ₂	Н	CF ₃	0	0
A468	C≡CCH ₂	CH₃OCH(CH₃)CH₂	Н	CF ₃	0	O
A469	C≡CCH ₂	CH ₃ OCH ₂ CH(CH ₃)	Н	CF ₃	0	0
A470	C≅CCH ₂	CH ₃ OCH ₂ C(CH ₃) ₂	Н	CF ₃	0	0
A471	C≡CCH ₂	CH₃OCH(CH₃)	Н	CF ₃	0	0
A472	C≡CCH ₂	CH ₃ OC(CH ₃) ₂	Н	CF ₃	0	0
A473	C≡CCH ₂	HC≡CCH ₂	Н	CF ₃	0	0
A474	C≡CCH ₂	H ₂ C=CHCH ₂	Н	CF ₃	0	0
A475	C≡CCH ₂	CH ₃ C≡CCH ₂	Н	CF ₃	0	0
A476	C≡CCH ₂	Сн	Н	CF ₃	0	0
A477	C≡CCH₂	ОСН	н	CF ₃	0	0
A478	C≡CCH ₂	ССН	Н	CF ₃	0	0

Radical	R ₁	R ₂	R ₄	R ₃	X ₁	р
A479	C≡CCH ₂	СН	Н	CF ₃	0	0
A480	C≡CCH₂	СН	Н	CF ₃	0	O
A481	C≡CCH ₂	СН	н	CF ₃	0	0
A482	C≡CCH ₂	СН	Н	CF ₃	0	0
A483	C≡CCH ₂	ОСН	Н	CF ₃	Ο.	0
A484	C≡CCH ₂	O_CH	Н	CF ₃	0	0
A485	C≡CCH ₂	ОСН	Н	CF ₃	0	0
A486	C≡CCH ₂	O CH	Н	CF₃	0	0
A487	C≡CCH ₂		Н	CF ₃	0	0
A488	C≡CCH ₂	OCH ₃	Н	CF ₃	0	0
A489	C≅CCH₂	ОН	Н	CF ₃	0	0
A490	C≡CCH ₂	OCH ₃	Н	CF ₃	0	0
A491	C≡CCH₂ ́		н	CF₃	0	0
A492	C≡CCH ₂	он	н	CF₃	Ο	0

Radical	R ₁	R ₂	R ₄	R ₃	X ₁	р
A493	C≡CCH ₂	CH ₃ CH ₃	Н	CF₃	0	0
A494	C≡CCH ₂	CH ₃	н	CF₃	. 0	0
A495	C≡CCH ₂	N N CH ₃	Н	CF ₃	O	0
A496	C≡CCH ₂		Н	CF ₃	0	0
A497	C≡CCH₂		. H	CF ₃	0	0
A498	C≡CCH ₂		Н	CF ₃	0	0
A499	C≡CCH₂	OCH ₃	Н	CF ₃	0	0
A500	C≡CCH ₂	OH N	Н	CF ₃	0	0
A501	C≡CCH ₂	OCH ₃	Н	CF ₃	0	0
A502	C≡CCH ₂	N OH	Н	CF ₃	0	0
A503	C≡CCH ₂	OCH ₃	н	CF ₃	0	0
A504	C≡CCH ₂	N OH	Н	CF₃	0	0

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Radical	R ₁	R ₂	R ₄	R ₃	X ₁	р
A505	C≡CCH ₂	C.N	Н	CF₃	0	0
A506	C≡CCH ₂	ON	Н	CF₃	0	0
A507	C≡CCH ₂	F OCH ₃	Н	CF ₃	. 0	. 0
A508	C≡CCH ₂	OCH ₃	Н	CF₃	0	0 ·
A509	C≡CCH ₂	CH=CH OCH ₃	Н	CF₃	0	O .
A510	C≡CCH ₂	CH ₂	Н	CF₃	O	0
A511	C≡CCH ₂	CH₂	н	CF ₃	0	0
A512	C≡CCH ₂	O_CH ₂	• н	CF₃	Ο	0
A513	C≡CCH ₂	CH ₂	Н	CF₃	0	0
A514	C≡CCH₂	O CH ₂	Н	CF₃	0	0
A515	C≡CCH ₂	CH ₂	Н	CF₃	0	0
A516	C≡CCH ₂	CH ₂	Н	CF ₃	Ο	0
A517	C≡CCH ₂	CH ₂	Н	CF₃	0	0
A518	C≡CCH₂	OCH ₂	Н	CF₃	0	0

Radical	R ₁	R ₂	R ₄	R ₃	X ₁	р
A519	C≡CCH₂	OCH ₂	Н	CF₃	0	0
A520	. C≡CCH ₂	OCH ₂	Н	CF₃	0	0
A521	C≡CCH ₂	O CH ₂	н	CF ₃	0	0
A522	C≡CCH ₂	CH ₂	Н	CF ₃	0	0
A523	C≡CCH₂	OCH ₃	н	CF ₃	0	0
A524	C≡CCH ₂	CH₂ OH	Н	CF ₃	0	0
A525	C≡CCH ₂	OCH ₃	Н	CF ₃	Ο	0
A526	C≡CCH ₂	CH ₂	Н	CF₃	0	0
A527	C≡CCH ₂	CH ₂	Н	CF₃	0	0
A528	C≡CCH ₂	CH ₃ OCH ₂ CH ₂	Н	CF ₃	0	0
A529	C≡CCH ₂	CH ₃ N N OCH₂CH₂ CH₃	Н	CF ₃	0	0
A530	C≡CCH₂	NNNOCH₂CH₂ CH₃	Н	CF ₃	0	0

Radical	R ₁	R ₂	R ₄	R ₃	X ₁	р
A531	C≡CCH ₂	CH ₂	Н	CF ₃	0	0
A532	C≡CCH ₂	CH ₂	Н	CF₃	O .	0
A533	C≡CCH ₂	CH ₂	н	CF ₃	O	0
A534	C≡CCH ₂	OCH ₃ CH ₂	Н	CF ₃	0	0
A535	C≡CCH ₂	OH CH₂	н	CF ₃	0	0
A536	C≡CCH ₂	OCH ₃ CH ₂	Н	CF ₃	0	0
A537	C≡CCH₂	OH CH₂	Н	CF ₃	Ο	0
A538	C≡CCH ₂	OCH ₃ CH ₂	н	CF ₃	0	Ò
A539	C≡CCH ₂	OH CH₂	н	CF ₃	0	0
A540	C≡CCH ₂	CH ₂	H	CF ₃	0	0
A541	C≡CCH ₂	CH ₂	н	CF₃	Ο	0
A542	C≡CCH₂	CH ₂ OCH ₃	Н	CF₃	0	0

Radical	R ₁	R ₂	R ₄	R ₃	X ₁	р
A543	C≡CCH ₂	OCH ₂ CH ₂ OCH ₃	Н	CF ₃	0	0
A544	C≡CCH ₂	OCH ₂ CH ₂	H	CF₃	Ο	0
A545	C≡CCH ₂	OCH ₂	Н	CF₃	0	0
A546	CH₂	CH₃	Н	CF ₂ CI	0	0
A547	CH ₂	CH₃CH₂	Н	CF ₂ CI	0	0
A548	CH ₂	(CH₃)₂CH	Н	CF ₂ Cl	0	0
A549	CH₂	PhCH ₂	Н	CF ₂ Cl	0	0
A550	CH ₂	CH ₃	Н	CF ₂ CI	S	0
A551	CH ₂	CH ₃	Н	CF ₂ CI	so	0
A552	CH ₂	CH ₃	H	CF ₂ CI	SO ₂	0
A553	CH ₂	CH ₃ CH ₂ CH ₂	Н	CF ₂ CI	Ο	0
A554	CH ₂	CH ₃ OCH ₂	Н	CF ₂ CI	0	0
A555	CH ₂	CH ₃ CH ₂ OCH ₂	Н	CF ₂ CI	0	0
A556	CH ₂	CH₃OCH₂CH₂	Н	CF ₂ CI	0	0
A557	CH ₂	CH3CH2OCH2CH2	Н	CF ₂ CI	0	O
A558	CH ₂	CH ₃ OC(CH ₃) ₂ CH ₂	Н	CF ₂ CI	0	0
A559	CH ₂	CH ₃ OCH(CH ₃)CH ₂	Н	CF ₂ CI	0	0
A560	CH₂	CH₃OCH₂CH(CH₃)	Н	CF ₂ CI	0	0
A561	CH₂	CH ₃ OCH ₂ C(CH ₃) ₂	Н	CF ₂ CI	0	0
A562	CH ₂	CH₃OCH(CH₃)	H	CF ₂ CI	0	0
A563	CH₂	CH ₃ OC(CH ₃) ₂	Н	CF ₂ CI	0	0
A564	CH ₂	HC≡CCH ₂	Н	CF ₂ CI	0	0
A565	CH ₂	H ₂ C=CHCH ₂	Н	CF ₂ Cl	0	0
A566	CH ₂	CH₃C≡CCH₂	Н	CF₂CI	0	0
A567	CH ₂	Сн	Н	CF ₂ CI	0	0.
A568	CH ₂	Ссн	Н	CF ₂ CI	0	0

Radical	R ₁	R ₂	R ₄ · R ₃	X ₁	р
A569	CH ₂	Ссн	H CF ₂ CI	. 0	0
A570	CH₂	СH ОДСН	H CF₂CI	0	0
A571	CH₂	СН	H CF₂CI	Ö	0
A572	CH₂	СН	H CF₂CI	0	0
A573	CH₂	CH	H CF₂CI	0	0
A574	CH₂	O_CH	H CF₂CI	0	0
A575	CH₂	O_CH	H CF₂CI	0	0
A576	CH₂	СН	H CF₂CI	0	0
A577	CH₂	O_CH	H CF₂CI	0	0
A578	CH₂		H CF₂CI	0	0
A579	. CH₂	CCH₃	H CF₂CI	0	0
A580	CH₂	ОН	H CF₂CI	0	0
A581	CH₂	OCH3	H CF₂CI	0	0
A582	CH₂	ОН	H CF₂CI	0	0 .

Radical	R ₁	R ₂	R ₄	R ₃	X ₁	p
A583	CH₂		Н	CF ₂ Cl	0	0
A584	CH₂	CH ₃ CH ₃	Н	CF₂CI	0	0
A585	CH₂	CH ₃	Н	CF₂CI	Ο	0
A586	CH₂	CH ₃	Н	CF₂CI	Ο	0
A587	CH₂		Н	CF₂CI	0	0
A588	CH₂		Н	CF₂CI	Ο	0
A589	CH ₂		Н	CF₂CI	0	0
A590	CH₂	OCH ₃	Н	CF₂CI	0	0
A591	CH₂	OH N	Н	CF₂CI	0	0
A592	CH₂	OCH ₃	Н	CF₂CI	0	0
A593	CH₂ .	OH	Н	CF₂CI	0	0
A594	CH₂	OCH ₃	н	CF₂CI	Ο	0

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Radical	R ₁	R ₂	R ₄	R ₃	X ₁	p
A595	CH₂	OH	Н	CF ₂ CI	0	0
A596	CH ₂	, CN	Н	CF₂CI	0	0
A597	CH₂	CN N	н	CF₂CI	O	0
A598	CH ₂	POCH3	Н	CF₂CI	0	0
A599	CH₂	OCH ₃	Н	CF₂CI	0	0
A600	CH₂	CH ₂ CH=CH	н	CF ₂ CI	0	0
A601	CH₂	CH ₂	Н	CF₂CI	Ο	0
A602	CH ₂	CH ₂	Н	CF₂CI	0	0
A603	CH ₂	OCH ₂	Н	CF₂CI	0	0
A604	CH ₂	CH ₂	Н	CF₂CI	0	0
A605	CH₂	O CH $_{\dot{z}}$	н	CF ₂ CI	0	0
A606	CH₂	CH ₂	Н	CF₂CI	Ο	0
A607	CH ₂	CH ₂	H	CF ₂ CI	0	0
A608	CH ₂	CH ₂	Н	CF₂CI	0	0

Radical	R ₁	R ₂	R ₄	R ₃	X ₁	р
A609	CH ₂	CH ₂	Н	CF ₂ CI	0	0
A610	CH₂	CH ₂	Н	CF₂CI	0	0
A611	CH₂	O CH ₂	Н	CF₂CI	0	0
A612	CH ₂	CH ₂	н	CF₂CI	0	0
A613	CH ₂	CH ₂	н	CF ₂ Cl	0	0
A614	CH₂	CH ₂ OCH ₃	Н	CF₂CI	0	0
A615	CH₂	OH OH	н	CF₂CI	0	0
A616	CH₂	CH ₂ OCH ₃	Н	CF ₂ CI	0	0
A617	CH₂	CH ₂	Н	CF₂CI	0	0
A618	CH₂	CH ₂	Н	CF₂CI	0	0
A619	CH₂	CH ₃ OCH ₂ CH ₂	Н	CF₂CI	0	0
A620	CH₂	CH ₃ OCH ₂ CH ₂	Н	CF₂CI	0	0

Radical	R ₁	R ₂	R ₄	R ₃	X ₁	р
A621	CH₂	CH ₃ OCH ₂ CH ₂	Н	CF ₂ Cl	0	0
A622	CH ₂	CH ₂	Н	CF ₂ CI	0	0
A623	CH₂	CH ₂	Н	CF₂CI .	O	0
A624	CH ₂	CH ₂	Н	CF ₂ Cl	0	0
A625	CH₂	OCH ₃ CH ₂	н	CF ₂ CI	0	0
A626	CH₂	OH CH ₂	Н	CF₂CI	0	0
A627	CH₂	OCH ₃	Н	CF₂CI	0	0
A628	CH₂	OH CH ₂	Н	CF₂CI	0	0
A629	CH₂	OCH ₃ CH ₂	Н	CF₂CI	0	0
A630	CH₂	OH CH₂	Н	CF₂CI	0	0
A631	CH₂	CH₂ ON	Н	CF₂CI	0	0
A632	CH₂	CH ₂	Н	CF₂CI	0	0

Radical	\mathbf{R}_1	R ₂	R ₄	R ₃	Х1	р
A633	CH₂	CH ₂ OCH ₃	Н	CF ₂ CI	0	0
A634	CH₂	OCH ₂ CH ₂	Н	CF₂CI		0
A635	CH₂	OCH ₂ CH ₂	Н	CF₂CI	0	0
A636	CH₂	OCH ₃	Н	CF₂CI	0	0
A637	CH ₂	CH ₃	Н	CHF ₂	0	0
A638	CH ₂	CH₂CH₃	Н	CHF ₂	0	0
A639	CH₂	(CH₃)₂CH	Н	CHF₂	0	0
A640	CH₂	PhCH₂	Н	CHF ₂	0	0
A641	CH₂	CH₃	Н	CHF ₂	S	0
A642	CH ₂	CH₃	Н	CHF ₂	0	0
A643	CH ₂	CH ₃	H	CHF ₂	0	0
A644	CH₂	CH₃OCH₂	Н	CHF ₂	О.	0
A645	CH₂	CH₃CH₂OCH₂	Н	CHF ₂	0	0
A646	CH ₂	CH ₃ OCH ₂ CH ₂	Н	CHF ₂	0	0
A647	CH₂	CH3CH2OCH2CH2	Н	CHF ₂	0	0
A648	CH₂	CH₃OC(CH₃)₂CH₂	Н	CHF ₂	0	0
A649	CH₂	CH₃OCH(CH₃)CH₂	Н	CHF ₂	0	0
A650	CH₂	CH ₃ OCH ₂ CH(CH ₃)	Н	CHF ₂	0	0
A651	CH₂	CH ₃ OCH ₂ C(CH ₃) ₂	Н	CHF ₂	0	0
A652	CH₂	CH₃OCH(CH₃)	Н	CHF ₂	0	0
A653	CH₂ .	CH ₃ OC(CH ₃)₂	Н	CHF ₂	0	0
A654	CH₂	HC≡CCH₂	Н	CHF ₂	0	0
A655	CH ₂	H ₂ C=CHCH ₂	Н	CHF ₂	0	0
A656	CH₂	CH₃C≡CCH₂	Н	CHF ₂	0	0
A657	CH ₂	Сн	Н	CHF ₂	0	0

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Radical	R ₁	R ₂	R ₄	R ₃	X ₁	р
A658	CH₂	Сн	Н	CHF ₂	0	0
A659	CH ₂	Ссн	Н	CHF₂	0	0
A660	CH ₂	СН	Н	CHF ₂	Ο	0
A661	CH ₂	Сн	Н	CHF ₂	0	0
A662	CH₂	СН	н	CHF ₂	0	0
A663	CH ₂	СН	Н	CHF ₂	0	0
A664	CH₂	СН	Н	CHF₂	0	0
A665	CH₂	O_CH	Н	CHF ₂	0	O
A666	CH ₂	ОСН	Н	CHF ₂	0	0
A667	CH ₂	O CH	Н	CHF ₂	0	0
A668	CH₂		Н	CHF ₂	0	o
A669	CH₂	OCH ₃	Н	CHF ₂	O	0
A670	CH₂	ОН	Н	CHF ₂	0	0
A671	CH₂ .	OCH ₃	н	CHF ₂	0	0

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Radical	R ₁	R ₂	R ₄	R ₃	X ₁	р
A672	CH₂		Н	CHF₂	0	0
A673	CH₂	óн	Н	CHF ₂	0	0
A674	CH ₂	CH ₃	н	CHF ₂	0	0
A675	CH₂	CH ₃	н	CHF₂	0	0
A676	CH₂	ĊH₃ N N CH₃	Н	CHF₂	Ο	0
A677	CH₂	SI.3	Н	CHF ₂	0	0
A678	CH ₂		Н	CHF ₂	0	0
A679	CH₂		Н	CHF ₂	0	0
A680	CH₂	OCH ₃	Н	CHF ₂	0	0
A681	CH₂	OH OH	Н	CHF ₂	0	0
A682	CH₂ .	OCH ₃	н	CHF ₂	S	0
A683	CH₂	N N N N N N N N N N N N N N N N N N N	н	CHF ₂	SO	0

Radical	R ₁	R ₂	R ₄	R ₃	X ₁	р
A684	CH₂	OCH ₃	Н	CHF₂	SO ₂	0
A685	CH₂	OH N	Н	CHF ₂	. 0	0
A686	CH₂		н	CHF₂	0	0
A687	CH ₂		н	CHF₂	0	0
A688	CH₂	F OCH ₃	Н	CHF ₂	0	0
A689	CH₂	OCH ₃	Н	CHF₂	0	0
A690	CH₂	CH=CH OCH ₃	Н	CHF ₂	0	0
A691	CH₂ .	CH ₂	Н	CHF ₂	0	0
A692	CH ₂	CH₂	Н	CHF ₂	0	0
A693	CH ₂	O CH ₂	. н	CHF ₂	0	0
A694	CH₂	CH₂	H	CHF ₂	0	0
A695	CH₂ .	O CH ₂	Н	CHF ₂	0	0
A696	CH₂	CH ₂	Н	CHF ₂	0	0
A697	CH ₂	CH ₂	Н	CHF₂	0	0

Radical	R ₁	R ₂	R ₄	R ₃	X ₁	р
A698	CH ₂	CH ₂	Н	CHF ₂	0	0
A699	CH ₂	O CH ₂	Н	CHF ₂	0	0
A700	CH ₂	CH ₂	н	CHF ₂	0	0
A701	CH₂	OCH ₂	Н	CHF ₂	0	0
A702	CH ₂	O CH ₂	Н	CHF ₂	0	0
A703	CH₂	CH ₂	Н	CHF ₂	0	0
A704	CH ₂	CH ₂ OCH ₃	Н	CHF ₂	0	0
A705	CH ₂	CH ₂	Н	CHF ₂	0	0
A706	CH ₂	CH ₂ OCH ₃	Н	CHF ₂	0	0
A707	CH ₂	CH ₂	Н	CHF₂	0	0
A708	CH₂	CH ₂	Н	CHF ₂	0	0
A709	CH ₂	CH ₃ OCH ₂ CH ₂	Н	CHF ₂	0	0

Radical	R ₁	R ₂	R ₄	R ₃	X ₁	р
A710	CH₂	CH ₃ N N OCH ₂ CH ₂	Н	CHF ₂	0	0
A711	CH₂	NNNOCH2CH2	Н	CHF ₂	0	
A712	CH₂	€ CH ₂	Н	CHF ₂	0	0
A713	CH₂	CH ₂	H	CHF ₂	0	0
A714	CH ₂	CH ₂	Н	CHF ₂	0	0
A715	CH₂	OCH ₃ CH ₂	Н	CHF₂	0	0
A716	CH₂	OH CH ₂	Н	CHF₂	0	0
A717	CH₂	OCH ₃	Н	CHF ₂	O	.0
A718	CH₂	OH CH ₂	Н	CHF₂	0	0
A719	CH₂	OCH ₃	н	CHF₂	0	. 0
A720	CH₂	OH CH ₂	Н	CHF ₂	0	0
A721	CH₂	CH ₂	Н	CHF ₂	0	0

Radical	R ₁	R ₂	R ₄	R ₃	X ₁	р
A722	CH₂	CH ₂	Н	CHF ₂	0	0
A723	CH ₂	CH ₂ OCH ₃	Н	CHF₂	0	0
A724	· CH ₂	OCH ₂ CH ₂	H	CHF₂	0	0
A725	CH₂	OCH ₂ CH ₂	H	CHF ₂	Ο	0
A726	CH ₂	OCH ₃	Н	CHF ₂	0	0
A727	CH ₂	CH ₃	Н	CF ₃	0	1
A728	CH ₂	CH₂CH₃	Н	CF ₃	0	1
A729	CH₂	(CH₃)₂CH	Н	CF ₃	0	1
A730	CH ₂	PhCH ₂	Н	CF ₃	0	1
A731	CH ₂	CH ₃	Н	CF ₃	S	1
A732	CH ₂	CH ₃	Н	CF ₃	so	1
A733	CH ₂	CH ₃	Н	CF ₃	SO ₂	1
A734	CH ₂	CH₃OCH₂ _.	Н	CF ₃	0	1
A735	CH ₂	CH ₃ CH ₂ OCH ₂	Н	CF ₃	Ο	1
A736	CH ₂	CH ₃ OCH ₂ CH ₂	Н	CF ₃	Ο	1
A737	CH ₂	CH₃CH₂OCH₂CH₂	Н	CF₃	0	1
A738	CH ₂	CH ₃ OC(CH ₃) ₂ CH ₂	Н	CF ₃	0	. 1
A739	CH ₂	CH₃OCH(CH₃)CH₂	Н	CF ₃	0	1
A740	CH ₂	CH₃OCH₂CH(CH₃)	Н	CF₃	O	1
A741	CH ₂	CH3OCH2C(CH3)2	Н	CF ₃	0	1
A742	CH ₂	CH₃OCH(CH₃)	Н	CF ₃	0	1
A743	CH ₂	CH ₃ OC(CH ₃) ₂	Н	CF ₃	0	1
A744	CH ₂	HC≡CCH ₂	Н	CF ₃	0	1
A745	CH ₂	H ₂ C=CHCH ₂	Н	CF ₃	0	1
A746	CH ₂	CH₃C≡CCH₂	Н	CF ₃	0	1

Radical	R ₁	R ₂	R ₄	R ₃	X ₁	р
A747	CH₂	Ссн	Н	CF ₃	0	1
A748	CH ₂	ОСН	н	CF ₃	0	1
A749	CH ₂	Сн	н	CF ₃	0	1
A750	CH ₂	СН	Н	CF₃	0	1
A751	· CH ₂	Сн	Н	CF ₃	0	1
A752	CH₂	CH CH	Н	CF ₃	0	1
A753	CH ₂	СН	Н	CF ₃	0	1
A754	CH ₂	O CH	Н	CF ₃	0	1
A755	CH ₂	ОСН	Н	CF₃	0	1
A756	CH₂	O CH	Н	CF ₃	0	1
A757	CH ₂	O_CH	Н	CF ₃	0	Ť
A758	CH₂		н	CF ₃	0	1
A759	CH₂	OCH ₃	Н	CF ₃	0	1
A760	CH₂ .	OH	Н	CF₃	0	1
A761	CH₂		Н	CF₃	Ο	1
		OCH ₃				

Radical	R ₁	R ₂	R ₄	R ₃	X ₁	р
A762	CH₂	ОН	Н	CF₃	0	1
A763	CH ₂	S	H	CF ₃	0	1
A764	CH₂	CH ₃ CH ₃	Н	CF ₃	0	1
A765	CH₂	CH ₃	Н	CF₃	0	
A766	CH₂	CH ₃	Н	CF ₃	0	. 1
A767	CH₂		Н	CF ₃	0	1
A768	CH ₂		Н	CF₃	0	1
A769	CH₂	N	Н	CF ₃	Ο	1
A770	CH₂	OCH ₃	н	CF ₃	0	1
A771	CH₂	OH OH	Н	CF ₃	0	. 1
A772	CH₂ ·	OCH ₃	Н	CF₃	0	1
A773	CH₂	N OH	Н	CF ₃	0	1
	·····		·			

Radical	R ₁	R ₂	R ₄	R ₃	Χ ₁	р
A774	CH₂	OCH ₃	Н	CF₃	0	1
A775	CH₂	OH N	Н	CF₃	0	1
A776	CH₂	C N	H	CF ₃	0	1
A777	CH₂		Н	CF ₃	. 0	1
A778	CH₂	POCH3	Н	CF ₃	0	1
A779	CH₂	OCH ₃	н	CF ₃	0	1
A780	CH₂	CH ₂ CH=CH	Н	CF₃	O	1
A781	CH₂ .	CH ₂	н	CF ₃	Ο	i
A782	CH ₂	CH ₂	н	CF ₃	0	1
A783	CH ₂	OCH ₂	н	CF ₃	0	1
A784	CH₂	CH ₂	Н	CF ₃	0	1
A785	CH₂ .	OCH ₂	Н	CF ₃	0	1
A786	CH ₂	CH₂	Н	CF ₃	0	1
A787	CH ₂	CH ₂	Н	CF ₃	0	1

Radical	R ₁	R ₂	R ₄	R ₃	X ₁	р
A788	CH₂	CH ₂	Н	CF ₃	0	1
A789	CH₂	CH ₂	Н	CF ₃	0	1
A790	CH₂	CH ₂	Н	CF ₃	O 79	1
A791	CH₂	OCH ₂	Н	CF₃	0	1
A792	CH₂	O CH ₂	Н	CF₃	0	1
A793	CH ₂	CH ₂	Н	CF ₃	0	1
A794	CH₂	CH ₂ OCH ₃	Н	CF ₃	0	1
A795	CH ₂	CH ₂	Н	CF ₃	0	1
A796	CH₂	CH ₂ OCH ₃	Н	CF₃	0	1
A797	CH₂	CH ₂	н	CF₃	0	1
A798	CH₂	CH ₂	Н	CF ₃	0	1
A799	CH₂	CH ₃ OCH ₂ CH ₂	Н	CF ₃	0	1

Radical	R ₁	R ₂	R ₄	R ₃	X ₁	р
A800	CH₂	CH ₃ N N OCH ₂ CH ₂	Н	CF₃	0	1
A801	CH₂	OCH ₂ CH ₂	Н	CF ₃	0	1
A802	CH₂	CH ₂	Н	CF ₃	0	. 1
A803	CH₂	CH ₂	Н	CF₃	. О	1
A804	CH₂	CH ₂	Ή	CF ₃	0	1
A805	CH₂	OCH ₃ CH ₂	Н	CF₃	O	1
A806	CH₂	OH CH ₂	Н	CF₃	0	1
A807	CH₂	OCH ₃ CH ₂	Н	CF ₃	0	1
A808	CH₂	OH CH ₂	Н	CF ₃	0	1
A809	CH₂	OCH ₃ CH ₂	Н	CF₃	0	1
A810	CH₂	OH CH₂	н	CF₃	O	1
A811	CH₂	CH ₂	Н	CF₃	0	1

Radical	R ₁	R ₂	R ₄	R ₃	X ₁	р
A812	CH₂	CH ₂	Н	CF₃	0	1
A813	CH₂	CH ₂ OCH ₃	Н	CF ₃	. O	1
A814	CH₂	OCH ₂ CH ₂	Н	CF ₃	0	1
A815	CH₂	OCH ₂ CH ₂	Н	CF₃	0	1
A816	CH₂	OCH ₂	Н	CF ₃	0	1
A817	CH₂	CH₃SCH₂CH₂	Н	CF ₃	0	0
A818	CH ₂	CH₃SOCH₂CH₂	Н	CF ₃	0	0
A819	CH₂	CH ₃ SO ₂ CH ₂ CH ₂	Н	CF ₃	0	0
A820	CH ₂	CH₃OCH₂CH₂	Н	CF ₂ CI	0	1
A821	CH ₂	CH ₃ OCH ₂ CH ₂	Н	CF₂H	0	1
A822	CH ₂	CH₃OCH₂CH₂	F	CF ₃	0	0
A823	CH ₂	CH₃ÒCH₂CH₂	CH₃	CF ₃	0	0
A824	CH₂	CH₃OCH₂CH₂	CH ₃	CF ₃	0	1
A825	CH ₂	CH ₃ OCH ₂ CH ₂	Н	CF ₃	S	0
A826	CH ₂	CH₃OCH₂CH₂	Н	CF ₃	so	0
A827	CH ₂	CH₃OCH₂CH₂	CH ₃	CF ₃	SO ₂	0
A828	CH ₂	CH ₃ SO ₂ CH ₂ CH ₂	CH ₃	CF ₃	0	0
A829	CH₂	CH3S N-N	H	CF ₃	S	0
A830	CH₂ ·	CH ₃ O N N OCH ₃	Н	CF ₃	S	0

Radical	R ₁	R ₂	R ₄	R ₃	X ₁	p
A831	CH₂	CH ₃ N CH ₃	CH₃	CF₃	S	0
A832	CH₂	N, H	CH ₃	CF₃	S	0
A833	CH ₂	CH₃C(O)	Н	CF₃	0	0
A834	CH ₂	CF₃CH₂	Н	CF ₃	0	0
A835	CH₂	CH3OCH2CH2OCH2CH2	Н	CF ₃	0	0
A836	CH₂	HC≡CCH₂CH₂	Н	CF ₃	0	0
A837	CH ₂	Сн	Н	CF₃	0	.0
A838	CH ₂	CH ₃ CH ₂ C(OCH ₃)HOCH ₂ CH ₂	Н	CF ₃	0	0
A839	CH ₂	(CH₃)₃CC(O)	Н	CF ₃	0	0
A840	CH ₂	CH₂≈CHCH₂OCH₂CH₂	Н	CF ₃	O	0
A841	CH ₂	CH ₃ CH ₂ CH ₂ OCH ₂ CH ₂	Н	CF₃	0	0
A842	CH ₂	CH ₂	Н	CF ₃	0	0
A843	CH₂	n-Heptyl-C(O)	Н	CF₃	0	0
A844	CH ₂	Phenyl-C(O)	Н	CF ₃	0	-0
A845	CH ₂	CF₃CH₂OCH₂CH₂	Н	CF ₃	0	0
A846	CH ₂	CH₃OCH₂CH₂CH₂	Н	CF ₃	0	0
A847	CH ₂	HOCH ₂ CH ₂ CH ₂	Н	CF ₃	0	0
A848	CH ₂	CH ₂	Н	CF₃	0	0
A849	CH₂	N≡CCH2CH2	Н	CF₃	0	0
A850	CH₂	CICH ₂ CH ₂	Н	CF ₃	0	0
A851	CH₂	ОСН	Н	CF ₃	0	0
A852	CH₂	OCH ₂	н	CF ₃	0	0
A853	CH₂	CH ₃ OCH ₂ C(Br)HCH ₂	Н	CF ₃	0	0

Radical	R ₁	R ₂	R ₄	R ₃	X ₁	р
A854	CH₂	€S CH ₂	H	CF₃	0	0
A855	CH₂	O_CH ₂	Н	CF₃	0	0
A856	CH ₂	HOCH ₂ CH ₂	Н	CF ₃	0	0
A857	CH ₂	O_O_CH ₂	Н	CF₃	0	0
A858	CH₂	CH ₃ (OCH ₂ CH ₂) ₃	Н	CF ₃	О	0
A859	· CH ₂	CH₃CH₂OC(CH₃)HOCH₂CH₂	Н	CF ₃	0	0
A860	CH ₂	n-Heptyl-C(O)OCH ₂ CH ₂	Н	CF ₃	0	0
A861	CH ₂	CH ₃ C(O)OCH ₂ CH ₂	н	CF ₃	0	0
A862	CH ₂	CH₃SO₂OCH₂CH₂	Н	CF ₃	0	0
A863	CH₂	O	H	CF₃	0	0
A864	CH₂	CH₃	Н	CF ₃	-NCH ₃ SO ₂ -	0
A865	CH ₂	HOCH₂C(OH)HCH₂	н	CF ₃	0	0
A866	CH ₂	Phenyl-C(O)OCH ₂ CH ₂	Н	CF ₃	. О	0
A867	CH ₂	t-Butyl-C(O)OCH ₂ CH ₂	Н	CF ₃	0	0
A868	CH ₂	CH₃OC(O)CH₂	Н	CF₃	0	0

In the table below, in the case of rings, the ring attachment points for the substituents A_1 and A_2 are at the carbon atom which is marked "C", for example

, in the case of open-chain structures, "
$$(CH_3)_2C$$
" denotes, for example,

In the formula A-Q, Q denotes Q₁

$$R_{21}$$
 R_{13}
 R_{22}
 A_{2}
 A_{1}
 O
 $(Q_{1}),$

and Q_1 denotes the following radicals B:

Radical	A ₁	A ₂	n	R ₂₁	R ₂₂	R ₁₃
B1	CH ₂	CH₂	0	ы	1.1	011
B2	CH ₂	_	0	H	Н	OH .
B3	CH ₂	CH₂	0	CH₃	Н	ОН
B4		CH ₂	0	CH₃	CH₃	ОН
B5	(CH ₃)CH	CH₂	0	CH₃	CH₃	OH.
B6	(CH ₃) ₂ C	CH₂	0	CH₃	CH₃	ОН
	CH	СН	0	CH₃	-	OH
B7	CH₂	CH₂	0	CH₃	CH ₂ =CHCH ₂	OH
B8	CH ₂	CH₂	0	CH₃	HC≡CCH ₂	ОН
B9	CH ₂	CH₂	0	CH ₃	CH₃S	ОН
B10	CH ₂	CH ₂	0	CH₃	CH₃SO	ОН
B11	CH ₂	CH₂	0	CH₃	CH₃SO₂	ОН
B12	CH₂	CH ₂	0	CH₃	CH₃O	ОН
B13	CH ₂	CH₂	0	CH₃	CH₃OC(O)	ОН
B14	CH₂	CH ₂	0	CH₃	CH₃CH₂OC(O)	ОН
B15	CH ₂	(CH₃)₂C	0	Н	Н	ОН
B16	Î	CH ₂	0	Н	Н	ОН
	\bigcirc c					
B17	 Cc	CH₂	0	Н	н	ОН
B18	Cc	CH ₂	0	CH₃	н	ОН
B19	Cc	CH₂	0	CH ₃	CH ₃	ОН
B20	Çc	CH ₂	0	Н	Н	ОН
B21	Сс	CH₂	0	CH₃	н	ОН
B22	Çc	CH₂	0	CH₃	CH₃	ОН
B23	(CH ₃) ₂ C	0	0	CH₃	CH₃	ОН
B24	CH ₂	0	0	CH₃	CH₃	ОН

Radica	l A ₁	A ₂	n	R ₂₁	R ₂₂	R ₁₃
B25	CH₃N	0	0	CH₃	CH₃	ОН
B26	<u></u>	0	0	CH ₃	CH ₃	ОН
B27	CH₃N	CH ₂	0	СН₃	CH₃	ОН
B28	CH ₃ N	(CH₃)CH	0	Н	Н	ОН
B29	CH₃N	(CH₃)CH	0	CH₃	н	ОН
B30	NH	(CH₃)C	0	Н	-	ОН
B31	NH	CH	0	CH₃	-	ОН
B32	CH₃N	(CH₃)C	0	Н	-	ОН
B33	CH₃N	CH	0	CH ₃	-	ОН
B34	0	(CH ₃) ₂ C	0	Н	-	ОН
B35	0	(CH ₃) ₂ C	0	CH ₃	СН₃	ОН
B36	0	(CH ₃) ₂ C	0	CH₃	н	ОН .
B37	0	(CH₃)C	0	Н	-	ОН
B38	0	CH	0	CH₃	-	ОН
B39	$(CH_3)_2C$	C=O	0	CH₃	CH₃	ОН
B40	(CH ₃) ₂ C	(OH)CH	0	СН₃	CH₃	ОН
B41	Cc	C=O	0	CH₃	CH₃	ОН
B42	 Cc	C=O	0	CH ₂	CH ₂	ОН
B43	(CH ₃) ₂ C	S-C	0	CH₃	CH₃	ОН
B44	(CH₃)₂C	CH ₃ O C	0	CH ₃	CH₃	ОН
B45	(CH ₃) ₂ C		0	CH₃	CH₃	ОН
B46	(CH ₃) ₂ C	Ç, c	0	СН₃	СН₃	ОН
B47	(CH ₃) ₂ C	HON=C	0	CH ₃	CH₃	ОН
B48	$(CH_3)_2C$	CH₃ON=C	0	CH₃	CH₃	ОН
B49	$(CH_3)_2C$	BnON≃C	0	СН₃	CH₃	ОН
B50	СН	0	1	Н	CH₂	ОН
B51	CH	C=O	1	Н	CH ₂	ОН

Radical	A ₁	A ₂	n	R ₂₁	R ₂₂	R ₁₃
B52	CH	CH ₂	1	Н	ĊH₂	ОН
B53	СН	CH₃N	1	Н	CH₂	ОН
B54	CH	CH₂CH₂	1	Н	CH ₂	ОН
B55	CH	C=O	2	Н	CH₂	ОН
B56	СН	CH ₂	2	Н	CH₂	ОН
B57	СН	CH ₂	1	Н	CH₂	ÇI
B58	СН	CH ₂	1	Н	CH₂	NH ₂
B59	СН	CH ₂	1	Н	CH₂	CH₃SO₂NH
B60	СН	CH ₂	1	Н	CH₂	CH₃OCH₂CH₂S
B61	СН	CH ₂	1	н	CH₂	CH₃OCH₂CH₂SO
B62	СН	CH ₂	1	Н	CH₂	CH3OCH2CH2SO2
B63	СН	CH ₂	1	Н	CH₂	(CH₃)₂NC(O)NH
B64	CH	CH ₂	1	Н	CH ₂	PhC(O)O
B65	CH	CH ₂	1	Н	CH₂	CH₃OC(O)O
B66	СН	CH ₂	1	Н	CH₂	CH₃(CH₂) ₇ S
B67	CH	CH ₂	1	н	CH₂	CH ₃ (CH ₂) ₇ SO
B68	СН	CH ₂	1	Н	CH₂	CH ₃ (CH ₂) ₇ SO ₂
B69	CH	CH₂	1	Н	CH₂	(CH ₃)₂NSO₂NH
B70	CH	CH ₂	1	Н	CH₂	PhS
B71	CH	CH₂	1	Н	CH₂	PhSO
B72	CH	CH ₂	1	Н	CH ₂	PhSO₂
B73	СН	CH₂	1	Н	CH₂	N S
B74	СН	CH ₂	1	н	CH₂	CH₃S—NNN S S
B75	$(CH_3)_2C$	C=O	0	CH ₃	CH₃	CI
B76	$(CH_3)_2C$	C=O	0	CH₃	CH₃	NH ₂
B77	$(CH_3)_2C$	C=O	0	CH ₃	CH₃	CH₃SO₂NH
B78	$(CH_3)_2C$	C=O	0	CH₃	CH₃	CH₃OCH₂CH₂S
B79	$(CH_3)_2C$	C=O	0	CH ₃	CH₃	CH₃OCH₂CH₂SO
B80	$(CH_3)_2C$	C=O	0	CH ₃	CH₃	CH3OCH2CH2SO2
B81	(CH ₃) ₂ C	C=O	0	CH ₃	CH₃	(CH ₃)₂NC(O)NH

Radical	A ₁	A ₂	'n	R ₂₁	R ₂₂	R ₁₃
B82	(CH ₃) ₂ C	C=O	0	CH ₃	CH ₃	PhC(O)O
B83	$(CH_3)_2C$	C=O	0	CH_3	CH ₃	CH ₃ OC(O)O
B84	$(CH_3)_2C$	C=O	0	CH ₃	CH₃	CH₃(CH₂) ₇ S
B85	$(CH_3)_2C$	C=O	0	CH ₃	CH₃	CH₃(CH₂) ₇ SO
B86	.(CH ₃) ₂ C	C=O	0	CH ₃	CH ₃	CH ₃ (CH ₂) ₇ SO ₂
B87	$(CH_3)_2C$	C=O	0	CH₃	CH₃	(CH ₃) ₂ NSO ₂ NH
B88	$(CH_3)_2C$	C=O	0	CH ₃	CH₃	PhS
B89	$(CH_3)_2C$	C=O	0	CH ₃	CH₃	PhSO
B90	$(CH_3)_2C$	C=O	0	CH ₃	CH₃	PhSO ₂
B91	$(CH_3)_2C$	C=O	0	CH ₃	CH ₃	N
						N s
B92	(CH₃)₂C	C=O	0	СН₃	CH₃	CH ₃ S— N N S S
B93	(CH ₃) ₂ C	CH ₂	0	Н	Н	CI
B94	$(CH_3)_2C$	CH ₂	0	Н	Н	NH ₂
B96	(CH ₃) ₂ C	CH ₂	0	Н	Н	CH₃OCH₂CH₂S
B97	(CH ₃)₂C	CH₂	0	Н	• .Н	CH₃OCH₂CH₂SO
B98	(CH ₃) ₂ C	CH ₂	0	Н	Н	CH ₃ OCH ₂ CH ₂ SO ₂
B99	(CH₃)₂C	CH ₂	0	Н	Н	(CH ₃) ₂ NC(O)NH
B100	(CH₃)₂C	CH ₂	0	Н	Н	PhC(O)O
B101	(CH ₃) ₂ C	CH₂	0	Н	Н	CH₃OC(O)O
B102	(CH ₃) ₂ C	CH ₂	0	Н	Н	CH₃(CH₂) ₇ S
B103	(CH₃)₂C	CH ₂	0	Н	Н	CH ₃ (CH ₂) ₇ SO
B104	$(CH_3)_2C$	CH ₂	0	Н	Н	CH ₃ (CH ₂) ₇ SO ₂
B105	$(CH_3)_2C$	CH ₂	0	Н	Н	(CH₃)₂NSO₂NH
B106	$(CH_3)_2C$	CH ₂	0	Н	Н	PhS
B107	$(CH_3)_2C$	CH ₂	0	Н	н	PhSO
B108	$(CH_3)_2C$	CH ₂	0	Н	H	PhSO₂
B109	(CH ₃) ₂ C	CH ₂	0	Н	Н	N. N.
						N S

Radical	A ₁	A ₂	n	R ₂₁	R ₂₂	R ₁₃
B110	(CH ₃) ₂ C	CH ₂	0	Н	Н	CH ₃ S
B111	CH ₂	(CH₃)CH	0	Н	Н	ОН
B112	CH ₂	CH ₂	1	Н	CH ₂	t-Butyl-C(O)O
B113	CH ₂	CH ₂	1	Н	CH ₂	t-Heptyl-C(O)O

or Q in the formula A-Q denotes \mathbf{Q}_2

and Q_2 denotes the following radicals C:

Radical	R ₃₄	R ₃₅	R ₃₆
C1	CH₃	Н	ОН
C2	CH₃	CH ₃	ОН
СЗ	Н	HC≡CCH ₂	ОН
C4	Н	CH₃SO₂	ОН
C5	Н	CH ₃	ОН
C6	Н	PhCH ₂	ОН
C7	CF ₃	CH ₃	ОН
C8	Ссн	CH ₃	ОН
C9	CH ₃ OCH ₂ CH ₂ OCH ₂	CH ₃	ОН
C10	H ·	CH ₃	CI
C11	Н	CH ₃	NH_2
C12	Н	CH₃	CH₃SO₂NH
C13	Н	CH₃	CH₃OCH₂CH₂S
C14	Н	CH₃	CH₃OCH₂CH₂SO
C15	Н	CH ₃	CH₃OCH₂CH₂SO₂

Radical	R ₃₄	R ₃₅	R ₃₆
C16	Н	CH₃	(CH₃)₂NC(O)NH
C17	Н	CH₃	PhC(O)O
C18	Н	CH₃	CH₃OC(O)O
C19	Н	CH₃	$CH_3(CH_2)_7S$
C20	Н	CH₃	CH ₃ (CH ₂) ₇ SO
C21	Н	CH₃	$CH_3(CH_2)_7SO_2$
C22	Н	CH ₃	(CH ₃) ₂ NSO ₂ NH
C23	н	CH ₃	PhS
C24	Н	CH ₃	PhSO
C25	Н	CH ₃	PhSO₂
C26	н	CH₃	N S
C27	Н	CH₃	CH ₃ S S
C28	Н	CH₃	CH₃SO₂O
C29	Н	CH₃	p-TolylSO₂O

or Q in the formula A-Q denotes Q₃

$$R_{50}$$
 S (Q_3) (Q_3)

and Q_3 denotes the following radicals D (the point of attachment of R_{49} to the heterocycle is the "CH" group):

Radical	R ₄₉	R ₅₀	n
D1	Сн	CH ₃	0

	Radical	R ₄₉	R ₅₀	n
•	D2	Сн	CH₃	1
	D3	Сн	CH₃	2
	D4	Сн	CF ₃	0
	D5	Сн	CF ₃	1
	D6	Сн	CF ₃	2
	D7	Сн	Ph	0
	D8	Сн	Ph	1
	D9	Сн	Ph	2
	D10	Сн	PhCH ₂	0
	D11	Сн	PhCH ₂	1
	D12	Сн	PhCH ₂	2

Table 1: Intermediates for preparing the compounds of the formula I, represented as formula

A-Q

in which Q denotes hydroxyl:

ОН	<u>OH</u>	<u>он</u>	<u>OH</u>	<u>OH</u>	ОН	OH	<u>OH</u>	<u>OH</u>	ОН	<u>OH</u>	ОН
-	-	-	-	-	-	-	A8	A9	A10	A11	A12
A13	A14	A15	A16	A17	A18	A19	A20	A21	A22	A23	A24
A25	A26	A27	A28	A29	A30	A31	A32	A33	A34	A35	A36
A37	A38	A39	A40	A41	A42	A43	A44	A45	A46	A47	A48
A49	A50	A51	A52	A53	A54	A55	A56	A57	A58	A59	A60
A61	A62	A63	A64	A65	A66	A67	A68	A69	A70	A71	A72
A73	A74	A75	A76	A77	A78	A79	08A	A81	A82	A83	A84

ОН	ОН	OH	ОН	<u>OH</u>	<u>OH</u>	<u>OH</u>	ОН	ОН	ОН	ОН	ОН
A85	A86	A87	A88	A89	A90	-	-	-	-		-
-	-	A99	A100	A101	A102	A103	A104	A105	A106	A107	A108
A109	A110	A111	A112	A113	A114	A115	A116	A117	A118	A119	A120
A121	A122	A123	A124	A125	A126	A127	A128	A129	A130	A131	A132
A133	A134	A135	A136	A137	A138	A139	A140	A141	A142	A143	A144
A145	A146	A147	A148	A149	A150	A151	A152	A153	A154	A155	A156
A157	A158	A159	A160	A161	A162	A163	A164	A165	A166	A167	A168
A169	A170	A171	A172	A173	A174	A175	A176	A177	A178	A179	A180
A181	A182	A183	A184	A185	A186	A187	A188	A189	A190	A191	A192
A193	A194	A195	A196	A197	A198	A199	A200	A201	A202	A203	A204
A205	A206	A207	A208	A209	A210	A211	A212	A213	A214	A215	A216
A217	A218	A219	A220	A221	A222	A223	A224	A225	A226	A227	A228
A229	A230	A231	A232	A233	A234	A235	A236	A237	A238	A239	A240
A241	A242	A243	A244	A245	A246	A247	A248	A249	A250	A251	A252
A253	A254	A255	A256	A257	A258	A259	A260	A261	A262	A263	A264
A265	A266	A267	A268	A269	A270	A271	A272	A273	A274	A275	A276
A277	A278	A279	A280	A281	A282	A283	A284	A285	A286	A287	A288
A289	A290	A291	A292	A293	A294	A295	A296	A297	A298	A299	A300
A301	A302	A303	A304	A305	A306	A307	A308	A309	A310	A311	A312
A313	A314	A315	A316	A317	A318	A319	A320	A321	A322	A323	A324
A325	A326	A327	A328	A329	A330	A331	A332	A333	A334	A335	A336
A337	A338	A339						A345			
A349								A357			
A361	A362	A363	A364	A365	A366	A367	A368	A369	A370	A371	A372
A373	A374	A375	A376	A377	A378	A379	A380	A381	A382	A383	A384
A385		A387			A390			A393	A394	A395	A396
A397					A402				A406	A407	A408
A409	A410				A414				A418	A419	A420
A421	A422	A423	A424	A425	A426	A427	A428	A429	A430	A431	A432
	A434				A438		A440	A441	A442	A443	A444
					A450			A453	A454	A455	A456
					A462				A466		A468
A469	A470	A471	A472	A473	A474	A475	A476	A477	A478	A479	A480

ОН	<u>OH</u>	<u>OH</u>	ОН	<u>OH</u>	ОН						
A481	A482	A483	A484	A485	A486	A487	A488	A489	A490	A491	A492
A493	8 A494	A495	A496	A497	A498	A499	A500	A501	A502	A503	A504
A505	A506	A507	A508	A509	A510	A511	A512	A513	A514	A515	A516
A517	A518	A519	A520	A521	A522	A523	A524	A525	A526	A527	A528
A529	A530	A531	A532	A533	A534	A535	A536	A537	A538	A539	A540
A541	A542	A543	A544	A545	-	-	-	-	-	-	-
-	A554	A555	A556	A557	A558	A559	A560	A561	A562	A563	A564
A565	A566	A567	A568	A569	A570	A571	A572	A573	A574	A575	A576
A577	A578	A579	A580	A581	A582	A583	A584	A585	A586	A587	A588
A589	A590	A591	A592	A593	A594	A595	A596	A597	A598	A599	A600
A601			A604								
			A616								
A625			A628								
-			-								
_			A652								
A661			A664								
A685			A688								
A697			A700								
A709			A712								
			A724								
			A736								
			A748								
			A760								
			A772								
			A784								
			A796								
			808A								
			A820	A821	A822	A823	A824	A825	A826	A827	A828
A829	A830	A831	A832	-	-	-	-	-	-	-	-

Table 2: Compounds of the formula I, represented as compounds of the formula

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 $\mbox{\sc A-Q}$ in which Q denotes Q1 and Q1 denotes the radical B52:

B52	<u>B52</u>	<u>B52</u>	<u>B52</u>	<u>B52</u>	<u>B52</u>	<u>B52</u>	B52	B52	B52	B52	B52
		·									
A1	A2	A3	A4	A5	A6	A7	A8	A9	A10	A11	A12
A13	A14	A15	A16	A17	A18	A19	A20	A21	A22	A23	A24
A25	A26	A27	A28	A29	A30	A31	A32	A33	A34	A35	A36
A37	A38	A39	A40	A41	A42	A43	A44	A45	A46	A47	A48
A49	A50	A51	A52	A53	A54	A55	A56	A57	A58	A59	A60
A61	A62	A63	A64	A65	A66	A67	A68	A69	A70	A71	A72
A73	A74	A75	A76	A77	A78	A79	A80	A81	A82	A83	A84
A85	A86	A87	A88	A89	A90	A91	A92	A93	A94	A95	A96
A97	A98	A99	A100	A101	A102	A103	A104	A105	A106	A107	A108
A109	A110	A111	A112	A113	A114	A115	A116	A117	A118	A119	A120
A121	A122	A123	A124	A125	A126	A127	A128	A129	A130	A131	A132
A133	A134	A135	A136	A137	A138	A139	A140	A141	A142	A143	A144
A145	A146	A147	A148	A149	A150	A151	A152	A153	A154	A155	A156
A157	A158	A159	A1.60	A161	A162	A163	A164	A165	A166	A167	A168
A169	A170	A171	A172	A173	A174	A175	A176	A177	A178	A179	A180
A181	A182	A183	A184	A185	A186	A187	A188	A189	A190	A191	A192
A193	A194	A195	A196	A197	A198	A199	A200	A201	A202	A203	A204
A205	A206	A207	A208	A209	A210	A211	A212	A213	A214	A215	A216
A217	A218	A219	A220	A221	A222	A223	A224	A225	A226	A227	A228
A229	A230	A231	A232	A233	A234	A235	A236	A237	A238	A239	A240
A241	A242	A243	A244	A245	A246	A247	A248	A249	A250	A251	A252
A253	A254	A255	A256	A257	A258	A259	A260	A261	A262	A263	A264
A265	A266	A267	A268	A269	A270	A271	A272	A273	A274	A275	A276
A277	A278	A279	A280	A281	A282	A283	A284	A285	A286	A287	A288
A289	A290	A291	A292	A293	A294	A295	A296	A297	A298	A299	A300
A301	A302	A303	A304	A305	A306	A307	A308	A309	A310	A311	A312
A313	A314	A315	A316	A317	A318	A319	A320	A321	A322	A323	A324

B52	B52	B52	B52	B52	<u>B52</u>	B52	B52	B52	B52	B52	B52
A325	5 A326	A327	7 A328	A329	A330	A331	A332	2 A333	3 A334	A335	A336
A337	7 A338	3 A339	A340	A341	A342	A343	A344	A345	A346	A347	' A348
A349	A350	A351	A352	A353	A354	A355	A356	A357	' A358	A359	A360
A361	A362	A363	3 A364	A365	A366	A367	A368	A369	A370	A371	A372
A373	3 A374	A375	A376	A377	A378	A379	A380	A381	A382	. A383	A384
A385	A386	A387	A388	A389	A390	A391	A392	A393	A394	A395	A396
A397	' A398	A399	A400	A401	A402	A403	A404	A405	A406	A407	A408
A409	A410	A411	A412	A413	A414	A415	A416	A417	A418	A419	A420
A421	A422	A423	A424	A425	A426	A427	A428	A429	A430	A431	A432
A433	A434	A435	A436	A437	A438	A439	A440	A441	A442	A443	A444
A445	A446	A447	A448	A449	A450	A451	A452	A453	A454	A455	A456
A457	A458	A459	A460	A461	A462	A463	A464	A465	A466	A467	A468
A469	A470	A471	A472	A473	A474	A475	A476	A477	A478	A479	A480
A481	A482	A483	A484	A485	A486	A487	A488	A489	A490	A491	A492
A493	A494	A495	A496	A497	A498	A499	A500	A501	A502	A503	A504
A505	A506	A507	A508	A509	A510	A511	A512	A513	A514	A515	A516
A517	A518	A519	A520	A521	A522	A523	A524	A525	A526	A527	A528
A529	A530	A531	A532	A533	A534	A535	A536	A537	A538	A539	A540
A541	A542	A543	A544	A545	A546	A547	A548	A549	A550	A551	A552
A553	A554	A555	A556	A557	A558	A559	A560	A561	A562	A563	A564
A565	A566	A567	A568	A569	A570	A571	A572	A573	A574	A575	A576
A577	A578	A579	A580	A581	A582	A583	A584	A585	A586	A587	A588
A589	A590	A591								A599	A600
A601									A610	A611	A612
A613			A616					A621	A622	A623	A624
A625			A628						A634		A636
A637			A640	A641	A642	A643	A644	A645	A646	A647	A648
A649		A651	A652					A657	A658	A659	A660
A661			A664						A670	A671	A672
A673			A676						A682	A683	A684
											A696
A697									A706		
A709	A710	A711	A712	A713	A714	A715	A716	A717	A718	A719	A720

<u>B52</u>	B52	B52	B52	<u>B52</u>	<u>B52</u>	B52	B52	<u>B52</u>	<u>B52</u>	B52	B52
A721	A722	A723	A724	A725	A726	A727	A728	A729	A730	A731	A732
A733	A734	A735	A736	A737	A738	A739	A740	A741	A742	A743	A744
A745	A746	A747	A748	A749	A750	A751	A752	A753	A754	A755	A756
A757	A758	A759	A760	A761	A762	A763	A764	A765	A766	A767	A768
A769	A770	A771	A772	A773	A774	A775	A776	A777	A778	A779	A780
A781	A782	A783	A784	A785	A786	A787	A788	A789	A790	A791	A792
A793	A794	A795	A796	A797	A798	A799	A800	A801	A802	A803	A804
A805	A806	A807	A808	A809	A810	A811	A812	A813	A814	A815	A816
A817	A818	A819	A820	A821	A822	A823	A824	A825	A826	A827	A828
A829	A830	A831	A832	-	-	-	-	-	-	-	-

Table 3: Compounds of the formula I, represented as compounds of the formula

 $\mbox{\sc A-Q}$ in which Q denotes $\mbox{\sc Q}_1$ and $\mbox{\sc Q}_1$ denotes the radical B39:

B39	<u>B39</u>	<u>B39</u>	<u>B39</u>	B39	<u>B39</u>	B39	B39	B39	B39	<u>B39</u>	B39
A1	A2	АЗ	A4	A5	A6	A7	8 A	A 9	A10	A11	A12
A13	A14	A15	A16	A17	A18	A19	A20	A21	A22	A23	A24
A25	A26	A27	A28	A29	A30	A31	A32	A33	A34	A35	A36
A37	A38	A39	A40	A41	A42	A43	A44	A45	A46	A47	A48
A49	A50	A51	A52	A53	A54	A55	A56	A57	A58	A59	A60
A61	A62	A63	A64	A65	A66	A67	A68	A69	A70	A71	A72
A73	A74	A75	A76	A77	A78	A79	A80	A81	A82	A83	A84
A85	A86	A87	A88	A89	A90	A91	A92	A93	A94	A95	A96
A97	A98	A99	A100	A101	A102	A103	A104	A105	A106	A107	A108
A109	A110	A111	A112	A113	A114	A115	A116	A117	A118	A119	A120
A121	A122	A123	A124	A125	A126	A127	A128	A129	A130	A131	A132
A133	A134	A135	A136	A137	A138	A139	A140	A141	A142	A143	A144
A145	A146	A147	A148	A149	A150	A151	A152	A153	A154	A155	A156

B39	B39	B39	B39	B39	B39	B39	B39	B39	B39	B39	B39
A157	' A158	A159	A160	A161	A162	A163					
A169	A170	A171	A172	A173	A174	A175	A176				
A181	A182	A183	3 A184	A185	A186	A187	A188	A189			A192
A193	A194	A195	A196	A197	A198	A199	A200	A201	A202		A204
A205	A206	A207	' A208	A209	A210	A211	A212	A213	A214	A215	A216
A217	A218	A219	A220	A221	A222	A223	A224	A225	A226	A227	A228
A229	A230	A231	A232	A233	A234	A235	A236	A237	A238	A239	A240
A241	A242	A243	A244	A245	A246	A247	A248	A249	A250	A251	A252
A253	A254	A255	A256	A257	A258	A259	A260	A261	A262	A263	A264
A265	A266	A267	A268	A269	A270	A271	A272	A273	A274	A275	A276
A277	A278	A279	A280	A281	A282	A283	A284	A285	A286	A287	A288
A289	A290	A291	A292	A293	A294	A295	A296	A297	A298	A299	A300
A301	A302	A303	A304	A305	A306	A307	A308	A309	A310	A311	A312
A313	A314	A315	A316	A317	A318	A319	A320	A321	A322	A323	A324
A325	A326	A327	A328	A329	A330	A331	A332	A333	A334	A335	A336
A337	A338	A339	A340	A341	A342	A343	A344	A345	A346	A347	A348
A349	A350	A351	A352	A353	A354	A355	A356	A357	A358	A359	A360
A361	A362	A363	A364	A365	A366	A367	A368	A369	A370	A371	A372
A373	A374	A375	A376	A377	A378	A379	A380	A381	A382	A383	A384
A385	A386	A387	A388	A389	A390	A391	A392	A393	A394	A395	A396
A397	A398	A399	A400	A401	A402	A403	A404	A405	A406	A407	A408
A409	A410	A411	A412	A413	A414	A415	A416	A417	A418	A419	A420
A421	A422			A425				A429	A430	A431	A432
			A436								
			A448								
			A460								
			A472								
A481			A484								
			A496								
			A508								
			A520								
			A532								
A541 	A542	A543	A544	A545	A546	A547	A548	A549	A550	A551	A552

B39	<u>B39</u>	B39	<u>B39</u>	<u>B39</u>	B39	<u>B39</u>	<u>B39</u>	<u>B39</u>	<u>B39</u>	<u>B39</u>	B39
A553	A554	A555	A556	A557	A558	A559	A560	A561	A562	A563	A564
A565	A566	A567	A568	A569	A570	A571	A572	A573	A574	A575	A576
A577	A578	A579	A580	A581	A582	A583	A584	A585	A586	A587	A588
A589	A590	A591	A592	A593	A594	A595	A596	A597	A598	A599	A600
A601	A602	A603	A604	A605	A606	A607	A608	A609	A610	A611	A612
A613	A614	A615	A616	A617	A618	A619	A620	A621	A622	A623	A624
A625	A626	A627	A628	A629	A630	A631	A632	A633	A634	A635	A636
A637	A638	A639	A640	A641	A642	A643	A644	A645	A646	A647	A648
A649	A650	A651	A652	A653	A654	A655	A656	A657	A658	A659	A660
A661	A662	A663	A664	A665	A666	A667	A668	A669	A670	A671	A672
A673	A674	A675	A676	A677	A678	A679	A680	A681	A682	A683	A684
A685	A686	A687	A688	A689	A690	A691	A692	A693	A694	A695	A696
A697	A698	A699	A700	A701	A702	A703	A704	A705	A706	A707	A708
A709	A710	A711	A712	A713	A714	A715	A716	A717	A718	A719	A720
A721	A722	A723	A724	A725	A726	A727	A728	A729	A730	A731	A732
A733	A734	A735	A736	A737	A738	A739	A740	A741	A742	A743	A744
A745	A746	A747	A748	A749	A750	A751	A752	A753	A754	A755	A756
A757	A758	A759	A760	A761	A762	A763	A764	A765	A766	A767	A768
A769	A770	A771	A772	A773	A774	A775	A776	A777	A778	A779	A780
A781	A782	A783	A784	A785	A786	A787	A788	A789	A790	A791	A792
A793	A794	A795	A796	A797	A798	A799	A800	A801	A802	A803	A804
A805	A806	A807	A808	A809	A810	A811	A812	A813	A814	A815	A816
A817	A818	A819	A820	A821	A822	A823	A824	A825	A826	A827	A828
A829	A830	A831	A832	-	-	-	-	-	-	-	~

Table 4: Compounds of the formula I, represented as compounds of the formula

A-Q

in which Q denotes Q_1 and Q_1 denotes the radical B3:

| <u>B3</u> |
|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|
| _ | | | | | | | | | | | |

B3	B3	B3	B3	B3	B3	<u>B3</u>	<u>B3</u>	<u>B3</u>	<u>B3</u>	B3	B3
-	-	_	_	-	_	_	A8	A9	A10	A11	A12
A13	A14	A15	A16	A17	A18	A19	A20		A22	A23	
A25	A26	A27	A28	A29	A30	A31	A32			A35	
A37	A38	A39	A40	A41	A42	A43	A44	A45	A46	A47	
A49	A50	A51	A52	A53	A54	A55	A56	A57	A58	A59	A60
A61	A62	A63	A64	A65	A66	A67	A68	A69	A70	A71	A72
A73	A74	A75	A76	A77	A78	A79	A80	A81	A82	A83	A84
A85	A86	A87	A88	A89	A90	_	-	_	-	-	-
-	-	A99	A100	A101	A102	A103	A104	A105	A106	A107	A108
A109	A110	A111	A112	A113	A114	A115	A116	A117	A118	A119	A120
A121	A122	A123	A124	A125	A126	A127	A128	A129	A130	A131	A132
A133	A134	A135	A136	A137	A138	A139	A140	A141	A142	A143	A144
A145	A146	A147	A148	A149	A150	A151	A152	A153	A154	A155	A156
A157	A158	A159	A160	A161	A162	A163	A164	A165	A166	A167	A168
A169	A170	A171	A172	A173	A174	A175	A176	A177	A178	A179	A180
A181	A182	A183	A184	A185	A186	A187	A188	A189	A190	A191	A192
A193	A194	A195	A196	A197	A198	A199	A200	A201	A202	A203	A204
A205	A206	A207	A208	A209	A210	A211	A212	A213	A214	A215	A216
A217	A218	A219	A220	A221	A222	A223	A224	A225	A226	A227	A228
A229	A230	A231	A232	A233	A234	A235	A236	A237	A238	A239	A240
A241	A242	A243	A244	A245	A246	A247	A248	A249	A250	A251	A252
A253	A254			A257					A262		A264
									A274		
									A286		
A289									A298		
									A310		
									A322		
									A334		
									A346	A347	A348
									A358	A359	A360
									A370		
A373	A374	A375	A376	A377	A378	A379	A380	A381	A382	A383	A384

<u>B3</u>	<u>B3</u>	<u>B3</u>	<u>B3</u>	<u>B3</u>	<u>B3</u>	<u>B3</u>	<u>B3</u>	<u>B3</u>	<u>B3</u>	<u>B3</u>	<u>B3</u>
A385	A386	A387	A388	A389	A390	A391	A392	A393	A394	A395	A396
A397	A398	A399	A400	A401	A402	A403	A404	A405	A406	A407	A408
A409	A410	A411	A412	A413	A414	A415	A416	A417	A418	A419	A420
A421	A422	A423	A424	A425	A426	A427	A428	A429	A430	A431	A432
A433	A434	A435	A436	A437	A438	A439	A440	A441	A442	A443	A444
A445	A446	A447	A448	A449	A450	A451	A452	A453	A454	A455	A456
A457	A458	A459	A460	A461	A462	A463	A464	A465	A466	A467	A468
A469	A470	A471	A472	A473	A474	A475	A476	A477	A478	A479	A480
A481	A482	A483	A484	A485	A486	A487	A488	A489	A490	A491	A492
A493	A494	A495	A496	A497	A498	A499	A500	A501	A502	A503	A504
A505	A506	A507	A508	A509	A510	A511	A512	A513	A514	A515	A516
A517	A518	A519	A520	A521	A522	A523	A524	A525	A526	A527	A528
A529	A530	A531	A532	A533	A534	A535	A536	A537	A538	A539	A540
A541	A542	A543	A544	A545	-	-	•	-	-	-	-
-					A558						
A565					A57.0						
A577					A582						
A589					A594						
					A606						
					A618						
A625	A626	A627	A628		A630						
* 0.40	-	-	-		-						
					A654						
					A666						
					A690						
					A702						
					A714						
		A723							-		
					A738						
					A750 A762						
					A774						
					A774						
	7.702	7/00	A/04	A/05	7/00	7/0/	W100	M/09	M/90	MIEIM	A/92

| <u>B3</u> |
|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|
| A793 | A794 | A795 | A796 | A797 | A798 | A799 | A800 | A801 | A802 | A803 | A804 |
| A805 | A806 | A807 | A808 | A809 | A810 | A811 | A812 | A813 | A814 | A815 | A816 |
| A817 | A818 | A819 | A820 | A821 | A822 | A823 | A824 | A825 | A826 | A827 | A828 |
| A829 | A830 | A831 | A832 | - | - | - | - | - | - | - | - |

Table 5: Compounds of the formula I, represented as compounds of the formula

 $\mbox{\sc A-Q}$ in which Q denotes $\mbox{\sc Q}_2$ and $\mbox{\sc Q}_2$ denotes the radical C5:

<u>C5</u>	<u>C5</u>	<u>C5</u>	<u>C5</u>	<u>C5</u>	<u>C5</u>	<u>C5</u>	<u>C5</u>	<u>C5</u>	<u>C5</u>	<u>C5</u>	<u>C5</u>
											
A1	A2	АЗ	A4	A 5	A6	A7	A8	A9	A10	A11	A12
A13	A14	A15	A16	A17	A18	A19	A20	A21	A22	A23	A24
A25	A26	A27	A28	A29	A30	A31	A32	A33	A34	A35	A36
A37	A38	A39	A40	A41	A42	A43	A44	A45	A46	A47	A48
A49	A50	A51	A52	A53	A54	A55	A56	A57	A58	A59	A60
A61	A62	A63	A64	A65	A66	A67	A68	A69	A70	A71	A72
A73	A74	A75	A76	A77	A78	A79	A80	A81	A82	A83	A84
A85	A86	A87	A88	A89	A90	A91	A92	A93	A94	A95	A96
A97	A98	A99	A100	A101	A102	A103	A104	A105	A106	A107	A108
A109	A110	A111	A112	A113	A114	A115	A116	A117	A118	A119	A120
A121	A122	A123	A124	A125	A126	A127	A128	A129	A130	A131	A132
A133	A134	A135	A136	A137	A138	A139	A140	A141	A142	A143	A144
A145	A146	A147	A148	A149	A150	A151	A152	A153	A154	A155	A156
A157	A158	A159	A160	A161	A162	A163	A164	A165	A166	A167	A168
A169	A170	A171	A172	A173	A174	A175	A176	A177	A178	A179	A180
A181	A182	A183	A184	A185	A186	A187	A188	A189	A190	A191	A192
A193	A194	A195	A196	A197	A198	A199	A200	A201	A202	A203	A204
A205	A206	A207	A208	A209	A210	A211	A212	A213		A215	A216
A217	A218	A219	A220	A221	A222	A223	A224			A227	A228

| <u>C5</u> |
|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|
| A229 | A230 | A231 | A232 | A233 | A234 | A235 | A236 | A237 | A238 | A239 | A240 |
| A241 | A242 | A243 | A244 | A245 | A246 | A247 | A248 | A249 | A250 | A251 | A252 |
| A253 | A254 | A255 | A256 | A257 | A258 | A259 | A260 | A261 | A262 | A263 | A264 |
| A265 | A266 | A267 | A268 | A269 | A270 | A271 | A272 | A273 | A274 | A275 | A276 |
| A277 | A278 | A279 | A280 | A281 | A282 | A283 | A284 | A285 | A286 | A287 | A288 |
| A289 | A290 | A291 | A292 | A293 | A294 | A295 | A296 | A297 | A298 | A299 | A300 |
| A301 | A302 | A303 | A304 | A305 | A306 | A307 | A308 | A309 | A310 | A311 | A312 |
| A313 | A314 | A315 | A316 | A317 | A318 | A319 | A320 | A321 | A322 | A323 | A324 |
| A325 | A326 | A327 | A328 | A329 | A330 | A331 | A332 | A333 | A334 | A335 | A336 |
| A337 | A338 | A339 | A340 | A341 | A342 | A343 | A344 | A345 | A346 | A347 | A348 |
| A349 | A350 | A351 | A352 | A353 | A354 | A355 | A356 | A357 | A358 | A359 | A360 |
| A361 | A362 | A363 | A364 | A365 | A366 | A367 | A368 | A369 | A370 | A371 | A372 |
| · A373 | A374 | A375 | A376 | A377 | A378 | A379 | A380 | A381 | A382 | A383 | A384 |
| A385 | A386 | A387 | A388 | A389 | A390 | A391 | A392 | A393 | A394 | A395 | A396 |
| A397 | A398 | A399 | A400 | A401 | A402 | A403 | A404 | A405 | A406 | A407 | A408 |
| A409 | A410 | A411 | A412 | A413 | A414 | A415 | A416 | A417 | A418 | A419 | A420 |
| A421 | A422 | A423 | A424 | A425 | A426 | A427 | A428 | A429 | A430 | A431 | A432 |
| A433 | A434 | A435 | A436 | A437 | A438 | A439 | A440 | A441 | A442 | A443 | A444 |
| A445 | A446 | A447 | A448 | A449 | A450 | A451 | A452 | A453 | A454 | A455 | A456 |
| A457 | A458 | A459 | A460 | A461 | | A463 | A464 | A465 | A466 | A467 | A468 |
| A469 | A470 | A471 | A472 | A473 | A474 | A475 | A476 | A477 | A478 | A479 | A480 |
| A481 | A482 | A483 | | A485 | | A487 | | A489 | A490 | A491 | A492 |
| A493 | A494 | A495 | | | | | | | A502 | | A504 |
| | | | | | | | | | A514 | A515 | A516 |
| | | A519 | | | | A523 | | | A526 | A527 | A528 |
| | A530 | A531 | | | | A535 | | | A538 | A539 | A540 |
| A541 | | A543 | | | | A547 | A548 | A549 | A550 | A551 | A552 |
| | | A555 | | A557 | A558 | A559 | A560 | A561 | A562 | A563 | A564 |
| | | A567 | | A569 | A570 | A571 | A572 | A573 | A574 | A575 | A576 |
| | | A579 | | A581 | A582 | A583 | A584 | A585 | A586 | A587 | A588 |
| A589 | | A591 | | A593 | | | A596 | A597 | A598 | A599 | A600 |
| A601 | | A603 | | | | A607 | | A609 | A610 | A611 | A612 |
| A613 | A614 | A615 | A616 | A617 | A618 | A619 | A620 | A621 | A622 | A623 | A624 |

| <u>C5</u> |
|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|
| A625 | A626 | A627 | A628 | A629 | A630 | A631 | A632 | A633 | A634 | A635 | A636 |
| A637 | A638 | A639 | A640 | A641 | A642 | A643 | A644 | A645 | A646 | A647 | A648 |
| A649 | A650 | A651 | A652 | A653 | A654 | A655 | A656 | A657 | A658 | A659 | A660 |
| A661 | A662 | A663 | A664 | A665 | A666 | A667 | A668 | A669 | A670 | A671 | A672 |
| A673 | A674 | A675 | A676 | A677 | A678 | A679 | A680 | A681 | A682 | A683 | A684 |
| A685 | A686 | A687 | A688 | A689 | A690 | A691 | A692 | A693 | A694 | A695 | A696 |
| A697 | A698 | A699 | A700 | A701 | A702 | A703 | Á704 | A705 | A706 | A707 | A708 |
| A709 | A710 | A711 | A712 | A713 | A714 | A715 | A716 | A717 | A718 | A719 | A720 |
| A721 | A722 | A723 | A724 | A725 | A726 | A727 | A728 | A729 | A730 | A731 | A732 |
| A733 | A734 | A735 | A736 | A737 | A738 | A739 | A740 | A741 | A742 | A743 | A744 |
| A745 | A746 | A747 | A748 | A749 | A750 | A751 | A752 | A753 | A754 | A755 | A756 |
| A757 | A758 | A759 | A760 | A761 | A762 | A763 | A764 | A765 | A766 | A767 | A768 |
| A769 | A770 | A771 | A772 | A773 | A774 | A775 | A776 | A777 | A778 | A779 | A780 |
| A781 | A782 | A783 | A784 | A785 | A786 | A787 | A788 | A789 | A790 | A791 | A792 |
| A793 | A794 | A795 | A796 | A797 | A798 | A799 | A800 | A801 | A802 | A803 | A804 |
| A805 | A806 | A807 | A808 | A809 | A810 | A811 | A812 | A813 | A814 | A815 | A816 |
| A817 | A818 | A819 | A820 | A821 | A822 | A823 | A824 | A825 | A826 | A827 | A828 |
| A829 | A830 | A831 | A832 | <u>-</u> | - | - | <u>.</u> | - | 7 | - | - |

Table 6: Compounds of the formula I, represented as compounds of the formula

 $\mbox{\bf A-Q}$ in which Q denotes $\mbox{\bf Q}_2$ and $\mbox{\bf Q}_2$ denotes the radical C2:

<u>C2</u>	<u>C2</u>	<u>C2</u>	<u>C2</u>								
			•								
A1	A2	АЗ	A4	A5	· A6	A7	A8	A 9	A10	A11	A12
A13	A14	A15	A16	A17	A18	A19	A20 .	A21	A22	A23	A24
A25	A26	A27	A28	A29	A30	A31	A32	A33	A34	A35	A36
A37	A38	A39	A40	A41	A42	A43	A44	A45	A46	A47	A48
									A58		

| <u>C2</u> |
|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|
| A61 | A62 | A63 | A64 | A65 | A66 | A67 | A68 | A69 | A70 | A71 | A72 |
| A73 | A74 | A75 | A76 | A77 | A78 | A79 | A80 | A81 | A82 | A83 | A84 |
| A85 | A86 | A87 | A88 | A89 | A90 | A91 | A92 | A93 | A94 | A95 | A96 |
| A97 | A98 | A99 | A100 | A101 | A102 | A103 | A104 | A105 | A106 | A107 | A108 |
| A109 | A110 | A111 | A112 | A113 | A114 | A115 | A116 | A117 | A118 | A119 | A120 |
| A121 | A122 | A123 | A124 | A125 | A126 | A127 | A128 | A129 | A130 | A131 | A132 |
| A133 | A134 | A135 | A136 | A137 | A138 | A139 | A140 | A141 | A142 | A143 | A144 |
| A145 | A146 | A147 | A148 | A149 | A150 | A151 | A152 | A153 | A154 | A155 | A156 |
| A157 | A158 | A159 | A160 | A161 | A162 | A163 | A164 | A165 | A166 | A167 | A168 |
| A169 | A170 | A171 | A172 | A173 | A174 | A175 | A176 | A177 | A178 | A179 | A180 |
| A181 | A182 | A183 | A184 | A185 | A186 | A187 | A188 | A189 | A190 | A191 | A192 |
| A193 | A194 | A195 | A196 | A197 | A198 | A199 | A200 | A201 | A202 | A203 | A204 |
| A205 | A206 | A207 | A208 | A209 | A210 | A211 | A212 | A213 | A214 | A215 | A216 |
| A217 | A218 | A219 | A220 | A221 | A222 | A223 | A224 | A225 | A226 | A227 | A228 |
| A229 | A230 | A231 | A232 | A233 | A234 | A235 | A236 | A237 | A238 | A239 | A240 |
| A241 | A242 | A243 | A244 | A245 | A246 | A247 | A248 | A249 | A250 | A251 | A252 |
| A253 | A254 | A255 | A256 | A257 | A258 | A259 | A260 | A261 | A262 | A263 | A264 |
| A265 | A266 | A267 | A268 | A269 | A270 | A271 | A272 | A273 | A274 | A275 | A276 |
| A277 | A278 | A279 | A280 | A281 | A282 | A283 | A284 | A285 | A286 | A287 | A288 |
| A289 | A290 | A291 | A292 | A293 | A294 | A295 | A296 | A297 | A298 | A299 | A300 |
| A301 | A302 | A303 | A304 | A305 | A306 | A307 | A308 | A309 | A310 | A311 | A312 |
| A313 | A314 | A315 | A316 | A317 | A318 | A319 | A320 | A321 | A322 | A323 | A324 |
| A325 | A326 | A327 | A328 | A329 | A330 | A331 | A332 | A333 | A334 | A335 | A336 |
| A337 | A338 | A339 | A340 | A341 | A342 | A343 | A344 | A345 | A346 | A347 | A348 |
| A349 | A350 | A351 | A352 | A353 | A354 | A355 | A356 | A357 | A358 | A359 | A360 |
| A361 | A362 | A363 | A364 | A365 | A366 | A367 | A368 | A369 | A370 | A371 | A372 |
| A373 | A374 | A375 | A376 | A377 | A378 | A379 | A380 | A381 | A382 | A383 | A384 |
| A385 | A386 | A387 | A388 | A389 | A390 | A391 | A392 | A393 | A394 | A395 | A396 |
| A397 | A398 | A399 | A400 | A401 | A402 | A403 | A404 | A405 | A406 | A407 | A408 |
| A409 | A410 | A411 | A412 | A413 | A414 | A415 | A416 | A417 | A418 | A419 | A420 |
| A421 | A422 | A423 | A424 | A425 | A426 | A427 | A428 | A429 | A430 | A431 | A432 |
| A433 | A434 | A435 | A436 | A437 | A438 | A439 | A440 | A441 | A442 | A443 | A444 |
| A445 | A446 | A447 | A448 | A449 | A450 | A451 | A452 | A453 | A454 | A455 | A456 |

	<u>C2</u>		<u>C2</u>		<u>C2</u>							
	A457	A458	A459	A460	A461	A462	A463	A464	A465	A466	A467	A468
	A469	A470	A471	A472	A473	A474	A475	A476	A477	A478	A479	A480
	A481	A482	A483	8 A484	A485	A486	A487	A488	A489	A490	A491	A492
	A493	A494	A495	A496	A497	A498	A499	A500	A501	A502	A503	A504
	A505	A506	A507	A508	A509	A510	A511	A512	A513	A514	A515	A516
	A517	A518	A519	A520	A521	A522	A523	A524	A525	A526	A527	A528
	A529	A530	A531	A532	A533	A534	A535	A536	A537	A538	A539	A540
	A541	A542	A543	A544	A545	A546	A547	A548	A549	A550	A551	A552
	A553	A554	A555	A556	A557	A558	A559	A560	A561	A562	A563	A564
	A565	A566	A567	A568	A569	A570	A571	A572	A573	A574	A575	A576
	A577	A578	A579	A580	A581	A582	A583	A584	A585	A586	A587	A588
	A589	A590	A591	A592	A593	A594	A595	A596	A597	A598	A599	A600
	A601	A602	A603	A604	A605	A606	A607	A608	A609	A610	A611	A612
	A613	A614	A615	A616	A617	A618	A619	A620	A621	A622	A623	A624
	A625	A626	A627	A628	A629	A630	A631	A632	A633	A634	A635	A636
	A637	A638	A639	A640	A641	A642	A643	A644	A645	A646	A647	A648
	A649	A650	A651	A652	A653	A654	A655	A656	A657	A658	A659	A660
	A661	A662	A663	A664	A665	A666	A667	A668	A669	A670	A671	A672
	A673	A674	A675	A676	A677	A678	A679	A680	A681	A682	A683	A684
	A685	A686	A687	A688	A689	A690	A691	A692	A693	A694	A695	A696
	A697	A698	A699	A700	A701	A702	A703	A704	A705	A706	A707	A708
	A709	A710	A711	A712	A713	A714	A715	A716	A717	A718	A719	A720
										A730	A731	A732
	A733	A734	A735	A736	A737	A738	A739	A740	A741	A742	A743	A744
	A745	A746	A747	A748	A749	A750	A751	A752.	A753	A754	A755	A756
	A757	A758	A759	A760	A761	A762	A763	A764	A765	A766	A767	A768
	A769	A770	A771	A772	A773	A774	A775	A776	A777	A778	A779	A780
	A781	A782	A783	A784	A785	A786	A787	A788	A789	A790	A791	A792
	A793	A794	A795	A796	A797	A798	A799	008A	A801	A802	A803	A804
	A805	A806	A807	808A	A809	A810	A811	A812	A813	A814	A815	A816
	A817	A818	A819	A820	A821	A822	A823	A824	A825	A826	A827	A828
	A829	A830	A831	A832	-	-	-	-	-	-	-	-
_												

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Table 7: Compounds of the formula I, represented as compounds of the formula

 $\mbox{\bf A-Q}$ in which Q denotes Q_2 and Q_2 denotes the radicals D1, D2 or D3:

	D1/D2/											
	<u>D3</u>											
	-	-	-	-	-	-	-	A8	A9	A10	A11	A12
	A13	A14	A15	A16	A17	A18	A19	A20	A21	A22	A23	A24
	A25	A26	A27	A28	A29	A30	A31	A32	A33	A34	A35	A36
	A37	A38	A39	A40	A41	A42	A43	A44	A45	A46	A47	A48
	A49	A50	A51	A52	A53	A54	A55	A56	A57	A58	A59	A60
	A61	A62	A63	A64	A65	A66	A67	A68	A69	A70	A71	A72
	A73	A74	A75	A76	A77	A78	A79	A80	A81	A82	A83	A84
	A85	A86	A87	A88	A89	A90	-	-	-	-	-	-
	-	-	A99	A100	A101	A102	A103	A104	A105	A106	A107	A108
	A109	A110	A111	A112	A113	A114	A115	A116	A117	A118	A119	A120
	A121	A122	A123	A124	A125	A126	A127	A128	A129	A130	A131	A132
	A133	A134	A135	A136	A137	A138	A139	A140	A141	A142	A143	A144
	A145	A146	A147	A148	A149	A150	A151	A152	A153	A154	A155	A156
	A157	A158	A159	A160	A161	A162	A163	A164	A165	A166	A167	A168
	A169	A170	A171	A172	A173	A174	A175	A176	A177	A178	A179	A180
	A181	A182	A183	A184	A185	A186	A187	A188	A189	A190	A191	A192
	A193	A194	A195	A196	A197	A198	A199	A200	A201	A202	A203	A204
	A205	A206	A207	A208	A209	A210	A211	A212	A213	·A214	A215	A216
	A217	A218	A219	A220	A221	A222	A223	A224	A225	A226	A227	A228
	A229	A230	A231	A232	A233	A234	A235	A236	A237	A238	A239	A240
	A241	A242	A243	A244	A245	A246	A247	A248	A249	A250	A251	A252
	A253	A254	A255	A256	A257	A258	A259	A260	A261	A262	A263	A264
	A265	A266	A267	A268	A269	A270	A271	A272	A273	A274	A275	A276
_	A277	A278	A279	A280	A281	A282	A283	A284	A285	A286	A287	A288

D1/D2/	D1/D2/	D1/D2/	D1/D2/	D1/D2/	D1/D2/	D1/D2/	D1/D2/	D1/D2/	D1/D2/	D1/D2/	D4/D0/
<u>D3</u>	D3	D3	<u>D3</u>	<u>D3</u>	<u>D3</u>	<u>D3</u>	<u>D3</u>	D3	D11021		
A289	A290	A291	A292	A293	A294	A295				<u>D3</u>	<u>D3</u>
A301	A302	A303	A304				A296	A297	A298	A299	A300
A313	A314			A305	A306	A307	A308	A309	A310	A311	A312
A325		A315	A316	A317	A318	A319	A320	A321	A322	A323	A324
	A326	A327	A328	A329	A330	A331	A332	A333	A334	A335	A336
A337	A338	A339	A340	A341	A342	A343	A344	A345	A346	A347	A348
A349	A350	A351	A352	A353	A354	A355	A356	A357	A358	A359	A360
A361	A362	A363	A364	A365	A366	A367	A368	A369	A370	A371	A372
A373	A374	A375	A376	A377	A378	A379	A380	A381	A382	A383	A384
A385	A386	A387	A388	A389	A390	A391	A392	A393	A394	A395	A396
A397	A398	A399	A400	A401	A402	A403	A404	A405	A406	A407	A408
A409	A410	A411	A412	A413	A414	A415	A416	A417	A418	A419	A420
A421	A422	A423	A424	A425	A426	A427	A428	A429	A430	A431	A432
A433	A434	A435	A436	A437	A438	A439	A440	A441	A442	A443	A444
A445	A446	A447	A448	A449	A450	A451	A452	A453	A454	A455	A456
A457	A458	A459	A460	A461	A462	A463	A464	A465	A466	A467	A468
A469	A470	A471	A472	A473	A474	A475	A476	A477	A478	A479	A480
A481	A482	A483	A484	A485	A486	A487	A488	A489	A490	A491	A492
A493	A494	A495	A496	A497	A498	A499	A500	A501	A502	A503	A504
A505	A506	A507	A508	A509	A510	A511	A512	A513	A514	A515	A516
A517	A518	A519	A520	A521	A522	A523	A524	A525	A526	A527	A528
A529	A530	A531	A532	A533	A534	A535	A536	A537	A538	A539	A540
A541	A542	A543	A544	A545	-	_	-	-	-	-	-
-	A554	A555	A556	A557	A558	A559	A560	A561	A562	A563	A564
A565	A566	A567	A568	A569	A570	A571	A572	A573	A574	A575	A576
A577	A578	A579	A580	A581	A582	A583	A584	A585	A586	A587	A588
A589	A590	A591	A592	A593	A594	A595	A596	A597	A598		A600
A601	A602	A603	A604	A605	A606	A607	A608	A609	A610		A612
A613	A614	A615	A616			A619		A621			A624
A625	A626					A631					A636
-	-	-	_	-	-	•					A648
A649	A650	A651	A652								A660
		· · · · · · · · · · · · · · · · · · ·	···							7008	A000

D3 D3<	D1/D2/	D1/D2/	D1/D2/	D1/D2/								
A685 A686 A687 A688 A689 A690 A691 A692 A693 A694 A695 A696 A697 A698 A699 A700 A701 A702 A703 A704 A705 A706 A707 A708 A709 A710 A711 A712 A713 A714 A715 A716 A717 A718 A719 A720 A721 A722 A723 A724 A725 A726 - <	<u>D3</u>	<u>D3</u>	<u>D3</u>	<u>D3</u>								
A697 A698 A699 A700 A701 A702 A703 A704 A705 A706 A707 A708 A709 A710 A711 A712 A713 A714 A715 A716 A717 A718 A719 A720 A721 A722 A723 A724 A725 A726 -	A661	A662	A663	A664	A665	A666	A667	A668	A669	A670	A671	A672
A709 A710 A711 A712 A713 A714 A715 A716 A717 A718 A719 A720 A721 A722 A723 A724 A725 A726 - <td>A685</td> <td>A686</td> <td>A687</td> <td>A688</td> <td>A689</td> <td>A690</td> <td>A691</td> <td>A692</td> <td>A693</td> <td>A694</td> <td>A695</td> <td>A696</td>	A685	A686	A687	A688	A689	A690	A691	A692	A693	A694	A695	A696
A721 A722 A723 A724 A725 A726 -	A697	A698	A699	A700	A701	A702	A703	A704	A705	A706	A707	A708
A733 A734 A735 A736 A737 A738 A739 A740 A741 A742 A743 A744 A745 A746 A747 A748 A749 A750 A751 A752 A753 A754 A755 A756 A757 A758 A759 A760 A761 A762 A763 A764 A765 A766 A767 A768 A769 A770 A771 A772 A773 A774 A775 A776 A777 A778 A779 A780 A781 A782 A783 A784 A785 A786 A787 A788 A789 A790 A791 A792 A793 A794 A795 A796 A797 A798 A799 A800 A801 A802 A803 A804 A805 A806 A807 A808 A809 A810 A811 A812 A813 A814 A815 A816	A709	A710	A711	A712	A713	A714	A715	A716	A717	A718	A719	A720
A745 A746 A747 A748 A749 A750 A751 A752 A753 A754 A755 A756 A757 A758 A759 A760 A761 A762 A763 A764 A765 A766 A767 A768 A769 A770 A771 A772 A773 A774 A775 A776 A777 A778 A779 A780 A781 A782 A783 A784 A785 A786 A787 A788 A789 A790 A791 A792 A793 A794 A795 A796 A797 A798 A799 A800 A801 A802 A803 A804 A805 A806 A807 A808 A809 A810 A811 A812 A813 A814 A815 A816	A721	A722	A723	A724	A725	A726	-	-	.	-	-	-
A757 A758 A759 A760 A761 A762 A763 A764 A765 A766 A767 A768 A769 A770 A771 A772 A773 A774 A775 A776 A777 A778 A779 A780 A781 A782 A783 A784 A785 A786 A787 A788 A789 A790 A791 A792 A793 A794 A795 A796 A797 A798 A799 A800 A801 A802 A803 A804 A805 A806 A807 A808 A809 A810 A811 A812 A813 A814 A815 A816	A733	A734	A735	A736	A737	A738	A739	A740	A741	A742	A743	A744
A769 A770 A771 A772 A773 A774 A775 A776 A777 A778 A779 A780 A781 A782 A783 A784 A785 A786 A787 A788 A789 A790 A791 A792 A793 A794 A795 A796 A797 A798 A799 A800 A801 A802 A803 A804 A805 A806 A807 A808 A809 A810 A811 A812 A813 A814 A815 A816	A745	A746	A747	A748	A749	A750	A751	A752	A753	A754	A755	A756
A781 A782 A783 A784 A785 A786 A787 A788 A789 A790 A791 A792 A793 A794 A795 A796 A797 A798 A799 A800 A801 A802 A803 A804 A805 A806 A807 A808 A809 A810 A811 A812 A813 A814 A815 A816	A757	A758	A759	A760	A761	A762	A763	A764	A765	A766	A767	A768
A793 A794 A795 A796 A797 A798 A799 A800 A801 A802 A803 A804 A805 A806 A807 A808 A809 A810 A811 A812 A813 A814 A815 A816	A769	A770	A771	A772	A773	A774	A775	A776	A777	A778	A779	A780
A805 A806 A807 A808 A809 A810 A811 A812 A813 A814 A815 A816	A781	A782	A783	A784	A785	A786	A787	A788	A789	A790	A791	A792
	A793	A794	A795	A796	A797	A798	A799	008A	A801	A802	A803	A804
A817 A818 A819 A820 A821 A822 A823 A824 A825 A826 A827 A828	A805	A806	A807	A808	A809	A810	A811	A812	A813	A814	A815	A816
	A817	A818	A819	A820	A821	A822	A823	A824	A825	A826	A827	A828
A829 A830 A831 A832	A829	A830	A831	A832	-	-	-	-	-	-	-	-

Table 8: Compounds of the formula lp:

in which R_1 , R_2 , R_3 , R_4 , X_1 and p have the same meaning as given for the radical A, and n is 0, 1 or 2:

A	A	A	A	A	A	A	A	A	<u>A</u>	A	<u>A</u>
											
-	-	-	-	-	-	-	A8	A9	A10	A11	A12
A13	A14	A15	A16	A17	A18	A19	A20	A21	A22	A23	A24
A25	A26	A27	A28	A29	A30	A31	A32	A33	A34	A35	A36
A37	A38	A39	A40	A41	A42	A43	A44	A45	A46	A47	A48
A49	A50	A51	A52	A53	A54	A55	A56	A57	A58	A59	A60
A61	A62	A63	A64	A65	A66	A67	A68	A69	A70	A71	A72
A73	A74	A75	A76	A77	A78	A79	A80	A81	A82	A83	A84
A85	A86	A87	A88	A89	A90	-	-	-	-	-	
-	-	A99	A100	A101	A102	A103	A104	A105	A106	A107	A108
A109	A110	A111	A112	A113	A114	A115	A116	A117	A118	A119	A120
A121	A122	A123	A124	A125	A126	A127	A128	A129	A130	A131	A132
A133	A134	A135	A136	A137	A138	A139	A140	A141	A142	A143	A144
A145	A146	A147	A148	A149	A150	A151	A152	A153	A154	A155	A156
A157	A158	A159	A160	A161	A162	A163	A164	A165	A166	A167	A168
A169	A170	A171	A172	A173	A174	A175	A176	A177	A178	A179	A180
A181	A182	A183	A184	A185	A186	A187	A188	A189	A190	A191	A192
A193	A194	A195	A196	A197	A198	A199	A200	A201	A202	A203	A204
		A207									
		A219									
		A231									
		A243									
	A254	A255	A256	A257	A258	A259	A260	A261	A262	A263	A264
A265	A266	A267		A269						A275	A276
A277	A278	A279	A280	A281	A282	A283	A284	A285	A286	A287	A288
A289	A290	A291	A292	A293	A294	A295	A296	A297	A298	A299	A300
A301		A303								A311	A312
A313	A314	A315	A316	A317	A318	A319	A320	A321	A322	A323	A324
A325	A326	A327	A328	A329	A330	A331	A332	A333	A334	A335	A336
A337	A338	A339	A340	A341	A342	A343	A344	A345	A346	A347	A348
A349	A350	A351	A352	A353	A354	A355	A356	A357	A358	A359	A360
A361	A362	A363	A364	A365	A366	A367	A368	A369	A370	A371	A372
A373	A374	A375	A376	A377	A378	A379	A380	A381	A382	A383	A384

							A		Α	Α	
<u>A</u>	<u>A</u>	<u>A</u>				<u>A</u>					
											A396
									A406		
A409	A410	A411	A412	A413	A414	A415	A416	A417	A418	A419	A420
A421	A422	A423	A424	A425	A426	A427	A428	A429	A430	A431	A432
A433	A434	A435	A436	A437	A438	A439	A440	A441	A442	A443	A444
A445	A446	A447	A448	A449	A450	A451	A452	A453	A454	A455	A456
A457	A458	A459	A460	A461	A462	A463	A464	A465	A466	A467	A468
A469	A470	A471	A472	A473	A474	A475	A476	A477	A478	A479	A480
A481	A482	A483	A484	A485	A486	A487	A488	A489	A490	A491	A492
A493	A494	A495	A496	A497	A498	A499	A500	A501	A502	A503	A504
A505	Ä506	A507	A508	A509	A510	A511	A512	A513	A514	A515	A516
A517	A518	A519	A520	A521	A522	A523	A524	A525	A526	A527	A528
A529	A530	A531	A532	A533	A534	A535	A536	A537	A538	A539	A540
A541	A542	A543	A544	A545	-	-	-	-	-	-	-
-	A554	A555	A556	A557	A558	A559	A560	A561	A562	A563	A564
A565	A566	A567	A568	A569	A570	A571	A572	A573	A574	A575	A576
A577	A578	A579	A580	A581	A582	A583	A584	A585	A586	A587	A588
A589	A590	A591	A592	A593	A594	A595	A596	A597	A598	A599	A600
A601	A602	A603	A604	A605	A606	A607	A608	A609	A610	A611	A612
A613	A614	A615	A616	A617	A618	A619	A620	A621	A622	A623	A624
A625	A626	A627	A628	A629	A630	A631	A632	A633	A634	A635	A636
-	-	-	-	•	-	-	A644	A645	A646	A647	A648
A649	A650	A651	A652	A653	A654	A655	A656	A657	A658	A659	A660
A661	A662	A663	A664	A665	A666	A667	A668	A669	A670	A671	A672
A685	A686	A687	A688	A689	A690	A691	A692	A693	A694	A695	A696
A697	A698	A699	A700	A701	A702	A703	A704	A705	A706	A707	A708
A709	A710	A711	A712	A713	A714	A715	A716	A717	A718	A719	A720
A721	A722	A723	A724	A725	A726	-	-	-	-	-	-
A733	A734	A735	A736	A737	A738	A739	A740	A741	A742	A743	A744
A745	A746	A747	A748	A749	A750	A751	A752	A753	A754	A755	A756
A757	A758	A759	A760	A761	A762	A763	A764	A765	A766	A767	A768
A769	A770	A771	A772	A773	A774	A775	A776	A777	A778	A779	A780
A781	A782	A783	A784	A785	A786	A787	A788	A789	A790	A791	A792

A	A	A	A	A	<u>A</u>	A	A	A	A	A	<u>A</u>
A793	A794	A795	A796	A797	A798	A799	A800	A801	A802	A803	A804
A805	A806	A807	A808	A809	A810	A811	A812	A813	A814	A815	A816
A817	A818	A819	A820	A821	A822	A823	A824	A825	A826	A827	A828
A829	A830	A831	A832	-	-	-	-	-	-	-	-

Table 9: Compounds of the formula I, represented as compounds of the formula

A-Q

in which A denotes A10:

<u>A10</u>	A10	<u>A10</u>	<u>A10</u>	<u>A10</u>	<u>A10</u>	<u>A10</u>	<u>A10</u>	A10	<u>A10</u>	<u>A10</u>	<u>A10</u>
									· · · · · · · · · · · · · · · · · · ·		
B1	B2	-	B4	B5	B6	B7	B8	B9	B10	B11	B12
B13	B14	B15	B16	B17	B18	B19	B20	B21	B22	B23	B24
B25	B26	B27	B28	B29	B30	B31	B32	B33	B34	B35	B36
B37	B38	-	B40	B41	B42	B43	B44	B45	B46	B47	B48
B49	B50	B51	-	B53	B54	B55	B56	B57	B58	B59	B60
B61	B62	B63	B64	B65	B66	B67	B68	B69	B70	B71	B72
B73	B74	B75	B76	B77	B78	B79	B80	B81	B82	B83	B84
B85	B86	B87	B88	B89	B90	B91	B92	B93	B94		B96
B97	B98	B99	B100	B101	B102	B103	B104	B105	B106	B107	B108
B109	B110	-	-	· -	-	-		•	-	-	-

Table 10: Compounds of the formula I, represented as compounds of the formula

A-Q

in which A denotes A10:

- 125 -

| <u>A10</u> |
|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|
| | | | | | | | | | | | |
| C1 | - | C3 | C4 | - | C6 | C7 | C8 | C9 | C10 | C11 | C12 |
| C13 | C14 | C15 | C16 | C17 | C18 | C19 | C20 | C21 | C22 | C23 | C24 |
| C25 | C26 | C27 | - | _ | - | - | - | _ | _ | - | |
| | | | | | | | | | | | |

Table 11: Compounds of the formula I, represented as compounds of the formula

A-Q

in which A denotes A10:

<u>A10</u>	A10	A10	<u>A10</u>								
_	-	•	D4	D5	D6	D7	D8	D9	D10	D11	D12

Table 12: Compounds of the formula I, represented as compounds of the formula

A-Q

in which A denotes A556:

A556	<u>A556</u>	<u>A556</u>	<u>A556</u>	<u>A556</u>	<u>A556</u>	<u>A556</u>	A556	A556	A556	<u>A556</u>	<u>A556</u>
					-				, ,,,,,,		
B1	B2	-	B4	B 5	B6.	B7	B8	B9	B10	B11	B12
B13	B14	B15	B16	B17	B18	B19	B20	B21	B22	B23	B24
B25	B26	B27	B28	B29	B30	B31	B32	B33	B34	B35	B36
B37	B38	-	B40	B41	B42	B43	B44	B45	B46	B47	B48
B49	B50	B51	-	B53	B54	B55	B56	B57	B58	B59	B60
B61	B62	B63	B64	B65	B66	B67	B68	B69	B70	B71	B72
B73	B74	B75	B76	B77	B78	B79	B80	B81	B82	B83	B84
B85	B86	B87	B88	B89	B90	B91	B92	B93	B94		B96
B97	B98	B99	B100	B101	B102	B103	B104	B105	B106	B107	B108

B109 B110	<u>A556</u>	<u>A556</u>	<u>A556</u>	<u>A556</u>	A556	A556	A556	A556	A556	<u>A556</u>	A556	A556
	B109	B110	-	-	-	-	-	-	-	-	-	-

Table 13: Compounds of the formula I, represented as compounds of the formula

A-Q

in which A denotes A556:

A556	<u>A556</u>	<u>A556</u>	<u>A556</u>	<u>A556</u>	<u>A556</u>	<u>A556</u> .	A556	<u>A556</u>	A556	A556	A556
C13	C14	C15	C16	C17	C18	C7 C19	C20	C21	C22		

Table 14: Compounds of the formula I, represented as compounds of the formula

A-Q

in which A denotes A556:

<u>A556</u>	A556	A556									
<u>.</u>	-	-	D4	D5	D6	D7	D8	D9	D10	D11	D12

Table 15: Compounds of the formula I, represented as compounds of the formula

A-Q

in which A denotes A646:

A646	<u>A646</u>	<u>A646</u>	A646	<u>A646</u>	A646	A646	A646	A646	<u>A646</u>	<u>A646</u>	<u>A646</u>
B1	B2	-	B4	B5	B6	B7	B8	B9	B10	B11	B12
B13	B14	B15	B16	B17	B18	B19	B20	B21	B22	B23	B24
B25	B26	B27	B28	B29	B30	B31	B32	B33	B34	B35	B36
B37	B38	-	B40	B41	B42	B43	B44	B45	B46	B47	B48
B49	B50	B51	-	B53	B54	B55	B56	B57	B58	B59	B60
B61	B62	B63	B64	B65	B66	B67	B68	B69	B70	B71	B72
B73	B74	B75	B76	B77	B78	B79	B80	B81	B82	B83	B84
B85	B86	B87	B88	B89	B90	B91	B92	B93	B94		B96
B97	B98	B99	B100	B101	B102	B103	B104	B105	B106	B107	B108
B109	B110	-	-	-	-	-	-	-	-	-	-

Table 16: Compounds of the formula I, represented as compounds of the formula

A-Q

in which A denotes A646:

| <u>A646</u> |
|-------------|-------------|-------------|-------------|-------------|-------------|-------------|-------------|-------------|-------------|-------------|-------------|
| C1 | - | C3 | C4 | - | C6 | C7 | C8 | C9 | C10 | C11 | C12 |
| | | | | C17 | | | | | | | • |
| C25 | C26 | C27 | - | - | - | - | - | - | - | - | _ |

Table 17: Compounds of the formula I, represented as compounds of the formula

A-Q

in which A denotes A646:

<u>A646</u>	A646	A646	<u>A646</u>	<u>A646</u>							
_	-	-	D4	D5	D6	D7	D8	D9	D10	D11	D12

Table 18: Physical data for the compounds of the formula I given in the tables above: (the melting points are given in °C.)

Compound	m.p. (range)	Phys. state
A1-C2	138-140	crystalline
A2-C2	138-140	crystalline
A833-B52 (K ⁺)	145-150	crystalline
A833-B52 (H4)	-	oil
A830-B52	-	amorphous/liquid
A829-B52	_	oil
A829-B1	-	oil
A10-B52 (H3)	54-56	crystalline
A10-B1	71-73	crystalline
A10-B3	-	viscous
A10-B14	-	viscous
A10-B39	99-100	crystalline
A736-B52	100-102	crystalline
A10-C2 (H6)	÷	viscous
A57-B52 (H5)	54-56	crystalline
A18-B52	71-74	crystalline
A8-B52	95-98	crystalline
A19-B52	53-55	crystalline
A1-C5	32-34	crystalline
A2-C5	32-33	crystalline
A10-C5	-	resin
A11-C5	38-39	crystalline
A11-B52	-	resin
A834-B52	-	crystalline
A835-B52	-	viscous
A854-B52	-	viscous
A90-B52	-	viscous

Compound	m.p. (range)	Phys. state
A33-B52	113-115	crystalline
A556-B52	-	crystalline
A646-B52	-	viscous
A868-B52	106-107	crystalline
A855-B52	-	viscous
A817-B52	-	viscous
A819-B52	-	crystalline
A856-B52	-	solid
A857-B52	-	viscous
A63-B52	-	resin
A20-B52	-	solid
A858-B52	-	resin
A836-B52	-	crystalline
A859-B52	-	viscous
A818-B52	-	viscous
A837-B52	-	viscous
A28-B52	-	viscous
A28-B52 (Et ₃ NH ⁺)	-	crystalline
A838-B52	-	viscous
A839-B52	- ·	viscous
A860-B52	-	viscous
A860-B113	-	viscous
A861-B52	90-93	crystalline
A840-B52	-	oil
A841-B52	41-43	crystalline
A842-B52	-	viscous
A843-B52	-	viscous
A866-B100	96-98	crystalline
A844-B52		viscous
A866-B112	-	viscous
A867-B112	-	viscous
A856-B112	79-81	crystalline
A20-C5	-	viscous
A10-C28	-	resin
A11-C28		resin

		
Compound	m.p. (range) Phys. state
A10-B52 (Et ₃ NH ⁺)	-	viscous
A862-B52	-	viscous
A24-B52	102-105	crystalline
A845-B52	40-44	crystalline
A837-B52	-	viscous
(Et₃NH⁺)		
A67-B52	68-69	crystalline
A863-B52	80-80	crystalline
A10-B17	40-42	crystalline
A846-B52	-	crystalline
A847-B52	-	viscous
A848-B52	-	crystalline
A56-B52	•	vitreous
A26-B52	-	vitreous
A849-B52	-	viscous
A10-B4	-	viscous
A865-B52	-	viscous
A850-B52	63-64	crystalline
A10-C29	•	resin
A10-B111	76-78	crystalline
A3-C5	-	resin
A834-C5	-	resin
A851-B52	-	vitreous
A852-B52	-	viscous
A10-B25	-	amorphous/liquid
A853-B52	-	viscous
A27-B52	-	oil
A864-C5	149-150	crystalline
A864-B52	110-112	crystalline
A834-B39	-	oil
A-852-OH	-	oil
A-851-OH	102-103	crystalline
A-835-OH	-	oil
A-24-OH	-	solid
A-858-OH	•	oil

Compound	m.p. (range)	Phys. state
A-859-OH	_	oil
A-864-OH	-	solid
A-851-OH	73-74	crystalline
A-848-OH	81-82	crystalline
A-27-OH	-	oil
A-855-OH	102-104	crystalline
A-90-OH	111-114	crystalline
A-124-OH	117-119	crystalline
A-834-OH	-	crystalline
A-852-OH	-	oil
A-851-OH	102-103	crystalline
A-835-OH	-	oil
A10-OH	62-63	crystalline
A830-OH	157-158	crystalline
A831-OH	188-189	crystalline
A829-OH	131-134	crystalline
A832-OH	110-112	crystalline

Table 19: Physical data for the compounds of the formula I given in the tables above: (the melting points given in °C.):

In the following formulas, end-standing valences denote methyl groups (in all cases except alkyne or alkene groups) or hydrogen (in the case of alkyne or alkene groups), for example

can be also drawn as
$$H_3C$$
 CH_3 and CH_3 and CH_3 CH_3

Comp.	Corresponding Formula	m.p. (range)
No.		Phys. state
1.001	F F O OH	138-140 crystalline
1.002	OH O ON F	145-150 crystalline
1.003	OH OF F	oil

o. (range) ys. state oil
oil
oil
oil
54-56 ystalline
ystalline
cous, 1H //R; 1.82 s); 3.26
3.37-3.39
n);3.57-
.60(m);
1(s); 4.84
7.74 (d); .82 (d)

Comp.	Corresponding Formula	m.p. (range)
No.	•	Phys. state
1.010	JUN OH SEF	viscous
1.011	N F F F	viscous
1.012	SH ST N	99-100 crystalline
1.013	OH OF PARTY OF	100-102 crystalline
1.014	P P P P P P P P P P P P P P P P P P P	viscous
1.015	F F F O O O O O O O O O O O O O O O O O	54-56 crystalline
1.016	OH OF PERSONS AND ADDRESS AND ADDRES	71-74 crystalline
1.017	OH OLON	95-98 crystalline

Comp.	Corresponding Formula	m.p. (range)
No.		Phys. state
1.018	OH OF P	53-55 crystalline
1.019	F F OH	32-34 crystalline
1.020	P P P P P P P P P P P P P P P P P P P	32-33 crystalline
1.021	F F F	resin
1.022	P F F N-	38-39 crystalline
1.023	OH O N F	resin
1.024	F F F F F F F F F F F F F F F F F F F	crystalline

Comp.	Corresponding Formula	m.p. (range)
No.		Phys. state
1.025	OHO CO ON F F	viscous
1.026	F F F O S	viscous
1.027	F F F OH	viscous
1.028	F F F OH	113-115 crystalline
1.029	OH O N F CI F	crystalline
1.030	OH O N F	viscous

Comp.	Corresponding Formula	m.p. (range) Phys. state
1.031	OH O N F F F	106-107 crystalline
1.032	F F F N O O O O O O O O O O O O O O O O	viscous
1.033	OH O ON F	viscous
1.034	OH O OFF	crystalline
1.035	OH O OH	crystalline
1.036	FFF ON OH OH	viscous

Comp.	Corresponding Formula	m.p. (range)
No.		Phys. state
1.037	F F F O O O O O O O O O O O O O O O O O	resin
1.038	OH O ON F	solid
1.039	OH O O O O O O O O O O O O O O O O O O	resin
1.040	OH O N F F F	crystalline
1.041	OHO ON F	viscous

Comp.	Corresponding Formula	m.p. (range)
No.		Phys. state
1.042	OH O N F F F	viscous
1.043	F F F O O O O O O O O O O O O O O O O O	viscous
1.044	F F F O O O O O O O O O O O O O O O O O	viscous
1.045	FTF OH	crystalline
1.046	OHO ON F	viscous
1.047	OH OF N F F	viscous

Comp.	Corresponding Formula	m.p. (range)
No.		Phys. state
1.048		viscous
1.049		viscous
1.050	OH O N F F F	90-93
1.051	OHO OF F	oil
1.052	OH O CO	41-43
1.053	F F F N O O O O O O O O O O O O O O O O	viscous

Comp.	Corresponding Formula	m.p. (range)
No.		Phys. state
1.054	OHO O OHO O OHO O FF	viscous
1.055		96-98
1.056	F F F O TO T	viscous
1.057	FFF OH	viscous
1.058	TO CONFE	viscous
1.059	OH N F F F	79-81

Comp.	Corresponding Formula	m.p. (range)
No.		Phys. state
1.060	O OH F N N N N N N N N N N N N N N N N N N N	viscous
1.061	F F F	viscous
1.062	F F F	resin
1.063	OHO CO	crystalline
1.064	OH O SO	viscous
1.065	F F F O CO	102-105 crystalline

Comp.	Corresponding Formula	m.p. (range)
No.		Phys. state
1.066	OHO	40-44 crystalline
1.067	FFF NO OH	viscous
1.068	F F F O CO O O O O O O O O O O O O O O O	68-69 crystalline
1.069	FFF ON OH	78-80 crystalline
1.070	AND PROPERTY OF THE PROPERTY O	40-42
1.071	° ° ° ° ° ° ° ° ° ° ° ° ° ° ° ° ° ° °	crystalline

Comp.	Corresponding Formula	m.p. (range) Phys. state
1.072	OH O I N F F	viscous
1.073	F F F N O O	46-47 crystalline
1.074	FFF N O O	vitreous
1.075	F F F	vitreous
1.076		75-76
1.077	N N N F F	viscous

Comp.	Corresponding Formula	m.p. (range)
NO.		Phys. state
1.078		viscous
1.079	OH OOH	viscous
1.080	CI N F F	63-64 crystalline
1.081	ONN ONN	resin
1.082	OH O N F F F	76-78 crystalline
1.083	O OH N N N N N N N N N N N N N N N N N N N	resin

Comp.	Corresponding Formula	m.p. (range)
No.		Phys. state
1.084	F F F	resin
1.085	F F F N O O O O O O O O O O O O O O O O	vitreous
1.086	F F F O O O O O O O O O O O O O O O O O	viscous
1.087	F O N.O OH	oil
1.088	OH O Br	viscous
1.089	F F F N O O O O O O O O O O O O O O O O	oil

Comp.	Corresponding Formula	m.p. (range)
No.		Phys. state
1.090	O:S- OH NF FF	149-150 crystalline
1.091	O OH N O OH	110-112 crystalline
1.092	OH FF	crystalline

Biological examples

Example B1: Herbicidal action before emergence of the plants (pre-emergence action)

Monocotyledonous and dicotyledonous test plants are sown in standard soil in pots. Immediately after sowing, the test substances are sprayed on at an optimum dosage (500 I of water/ha) as an aqueous suspension (prepared from a wettable powder (example F3, b) according to WO 97/34485) or emulsion (prepared from an emulsion concentrate (example F1, c) according to WO 97/34485). The test plants are then grown under optimum conditions in a greenhouse.

After a test period of 4 weeks, the test is evaluated using a 9-level scale of ratings (1 = complete damage, 9 = no effect). Ratings of 1 to 4 (in particular 1 to 3) mean good to very good herbicidal action.

Table B1: Pre-emergence action: ("NT" means "not tested"):

Compound	g/ha	Panicum	Digitaria	Echino.	Abutilon	Amaranthus	Chenop.
A10-B1	250	2	2	2	1	1	1
A10-B52, (H3)	250	1	1	1	1	.1	1
A830-B52	250	4	9	3	5	4	4
A1-C2	250	6	3	3	4	3	1
A833-B52 (K+)	250	1	2	2	1	2	1
A833-B52, (H4)	250	1	1	1 .	1	1	1
A10-B1	250	2	2	2	1	1	1
A10-B3	250	1	1	1	1	1	1
A10-B14	250	3	6	3	1	1	1
A10-B39	250	1	1	1	1	1	1
A736-B52	250	1	4	2	1	1	1
A10-C2 (H6)	250	3	3	3	1	2	1
A57-B52 (H5)	250	1	1	1	1	1	1
A18-B52	250	1	1	1	2	2	NT
A8-B52	250	1	1	1	1	1	NT
A19-B52	250	1	1	1	1	2	NT
A1-C5	250	2	2	1	2	2	1
A2-C5	250	1	2	2	1	1	1
A10-C5	250	2	3	1	1	1	1
A11-C5	250	1	2	1	1	1	1
A11-B52	250	1	1	1	1	2	1
A834-B52	250	1	1	1	1	2	1
A835-B52	250	1	2	1	2	1	2
A556-B52	250	1	1	1	1	2	1
A646-B52	250	1	1	1	1	2	1
A819-B52	250	7	9	7	1	2	1
A63-B52	250	2	3	1	5	3	NT
A20-B52	250	1	1	1	3	3	NT
A836-B52	250	1	2	1	5	2	3
A837-B52	250	1	2	2	1	2	NT
A28-B52	250	1	2	2	2	3	NT
A28-B52 (Et3NH+)	250	1	2	2	3	1	NT

					~~~~			·
	Compound	g/ha	Panicum	Digitaria	Echino.	Abutilon	Amaranthus	Chenop.
	A838-B52	250	1	3	2	1	1	1
	A839-B52	250	1	3	2	1	1	1
	A840-B52	250	1	2	2	2	2	1
•	A841-B52	250	1	2	1	1	1	1
	A842-B52	250	1	6	2	2	2	1
	A843-B52	250	· 1	2	2	1	1	1
	A844-B52	250	1	2	2	1	1	1
	A20-C5	250	1	2	2	1	1 .	1
	A10-C28	250	1	3	2	1	1	1
	A11-C28	250	2	2	1	2	1	1
	A10-B52 (Et3NH+)	250	1	1	2	1	1	1
·	A24-B52	250	.1	1	1	1	1	1
	A845-B52	250	1	1	1 ·	1	1	1
	A837-B52 (Et3NH+)	250	1	1	2	1	1	1
	A67-B52	250	1	2	2	3	2	1
	A10-B17	250	1	1	1	4	2	1
	A846-B52	250	1	1	1	2 .	1	1
	A847-B52	250	1	3	2	4	1	4
	A848-B52	250	1	1	1	7	1	1
	A56-B52	250	1	2	1	3	1	1 .
	A26-B52	250	1	1	1	1	1	1
	A849-B52	250	1	2	2	2	1	1
	A10-B4	250	2	3	1	3	1	1
	A850-B52	250	1	2	1	1	2	1
	A10-C29	250	2	2	1	1	1	NT
	A10-B111	250	1	1	1.	1	.1	NT
	A3-C5	250	1	2	2	1	1	NT ·
	A834-C5	250	1	3	1	1	2.	NT
	A851-B52 .	250	1	1	1 .	1	1	1
	A852-B52	250	1	1	1	4	1	2
	A10-B25	250	1	1	2	1	1	1
	A853-B52	250	1	1	2	1	1	2
	A27-B52	250	1	2	3	4	1	3

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The same results are obtained when the compounds of the formula I are formulated according to the other examples of WO 97/34485.

Example B2: Post-emergence herbicidal action

Monocotyledonous and dicotyledonous test plants are grown in standard soil in pots. At the 2- to 3-leaf stage of the test plants, the test substances are sprayed at optimum dosage (500 l of water/ha) as an aqueous suspension (prepared from a wettable powder (example F3, b) according to WO 97/34485) or emulsion (prepared from an emulsion concentrate (example F1, c) according to WO 97/34485). The test plants are then grown further under optimum conditions in a greenhouse.

After a test period of 2 to 3 weeks, the test is evaluated using a 9-level scale of rating (1 = complete damage, 9 = no effect). Ratings of 1 to 4 (in particular 1 to 3) mean good to very good herbicidal action.

Table B2: Post-emergence action:

	- , , - , , , ,								
Compound	g/ha	Panicum	Digitaria	Echino.	Abutilon	Xanth.	lpopur.	Amaranth	Chenop.
A10-B1	250	2	. 2	2	2	2	2	2	1
A10-B52, (H3)	250	1	1	2	1	2	2	2	1
A830-B52	250	4	9	3	5	4	5	4	· 4
A829-B52	250	2	6	4	3	6	4	2	2
A829-B1	250	7	9	7	7	4	6	2	2
A1-C2	250	7	8	4	3	4	3	2	4
A833-B52 (K+)	250	3	3	4	3	2	. 3	3	3
A833-B52, (H4)	250	3	3	4	3	1	2	2	3
A10-B3	250	1	1	3	1	1	2	1	1 .
A10-B14	250	2	2	3	2	2	2	2	1
A10-B39	250	. 1	3	3	1	2	2	1	1
A736-B52	250	1	1	2	1	2	2	1	1
A10-C2 (H6)	250	2	4	3	2	1	3	1	1
A57-B52 (H5)	250	1	1	2	1	2	2	1	1
A18-B52	250	1	1	2	1	2	2	1	1
A8-B52	250	1	1	1	1	2	1	1	1
A19-B52	250	1	1	2	1	2	1	1	1

Compound	g/ha f	anicum	Digitaria	Echino.	Abutilon	Xanth.	lpopur.	Amaranth	Chenop.
A1-C5	250	4	2	2	2	4	2	2	1
A2-C5	250	1	2	2	2	2	2	1	1
A10-C5	250	2	2	2	2	2	2	1	1
A11-C5	250	1	2	2	2	2	2	2	1
A11-B52	250	1	1	2	2	2	2	1	1
A834-B52	250	1	1 .	: 2	2	2	2	1	1
A835-B52	250	1	1	2	2	3	2	2	1
A854-B52	250	1	2	2	2	2	2	1	1
A90-B52	250	2	2	3	2	3	4	3	1
A33-B52	250	2	2	2	2	3	2	2	1
A556-B52	250	1	2	2	1	2	2	2	1
A646-B52	250	1	2	2	2	2	2	2	1
A855-B52	250	2	2	2	2	2	2	2	• 1
A817-B52	250	1	2	2	2	2	2	2	1
A819-B52	250	2	3	2	2	2	2	2	1
A856-B52	250	2	2	2	2	2	1	2	1
A857-B52	250	2	2	2	2	2	2	2	1
A63-B52	250	1	2	2	2	2	2	2	2
A20-B52	250	1	2	2	2	2	2	2	1
A858-B52	250	2	2	2	2	2	2	1	2
A836-B52	250	2	2	2	2	3	3	2	- 1
A859-B52	250	1	2	2	2	2	2	2	1
A818-B52	250	2	2	3	2	2	2	2	1
A837-B52	250	1	2	2	2	2	2	1	1
A28-B52	250	1	2	2	2	3	4	1	1
A28-B52									
(Et3NH+)	250	1	2	2	2	3	2	2	1
A838-B52	250	2	2	3	1	2	2	2	1
A839-B52	250	. 2	2	2	2	2	2	2	1
A860-B52	250	2	1	2	2	2	2	2	1
A861-B52	250	2	3	5	3 .	2	2	2	1
A840-B52	250	2	3	4	3	3	3	2	1
A841-B52	250	2	4	4	3	3	3	1	1
A842-B52	250	3	3	5	3	3	3	2	1
A843-B52	250	2	3	3	3	3	6	3	1

Compound	g/ha	Panicum	Digitaria	Echino.	Abutilon	Xanth.	lpopur.	Amaranth	Chenop.
A844-B52	250	2	3	4	3	3	3	3	1
A856-B112	250	3	3	5	2	3	3	3	1
A20-C5	250	2	3	4	3	3	3	2	1
A10-C28	250	4	4	4	3	3	3	2	1
A11-C28	250	3	4	4	3	3	3	2	1
A10-B52									
(Et3NH+)	250	2	2	2	2	2	2	2	1
A862-B52	250	2	2	3	3	2	5	2	1
A24-B52	250	2	2	2	2	2	2	2	1
A845-B52	250	2	2	2	2	2	2	2	1
A837-B52						•			
(Et3NH+)	250	2	2	2	2	2	2	2	1
A67-B52	250	2	2	2	2	2	3	2	1
A863-B52	250	2	2	3	2	2	3	2	1
A10-B17	250	2	1	2	2	2	. 2	2	1
A846-B52	250	1	2	2	2	2	1	2	2
A847-B52	250	1	2	1	2	2	4	3	1
A848-B52	250	2	2	2	2	2	2	1	1
A56-B52	250	1	2	2	2	2	2	1	1
A26-B52	250	2	2	2	2	2	2	2	1
A849-B52	250	2	2	3	2	2	2	2	2
A10-B4	250	1 .	1	2	1	2	1	2	1
A850-B52	250	2	2	2	2	2	2	2	1
A10-C29	250	2	3	3	2	2	1	2	1 ·
A10-B111	250	2	2	3	3	3	3	2	1
A3-C5	250	3	4	3	3	3	3	3	1
A851-B52	250	3	3	4	3	3	3	2	1 '
A852-B52	250	3	4	4	3	3	3	3	1
A10-B25	250	.3	4	4	3	3	3	2	1
A27-B52	250	1	2	2	3	2	4	2	5
4864-C5	250	1	2	2	2	2	2	1	1
A864-B52	250	2	2	2	2	2	2	2	1

The same results are obtained when the compounds of the formula I are formulated according to the other examples of WO 97/34485.

:1

WHAT IS CLAIMED IS:

1. A compound of the formula I

in which

p is 0 or 1;

 R_1 is a C_1 - C_6 alkylene, C_3 - C_6 alkenylene or C_3 - C_6 alkynylene chain which may be mono- or polysubstituted by halogen or R_5 , where the unsaturated bonds of the chain are not attached directly to the substituent X_1 ;

 X_1 is oxygen, -O(CO)-, -(CO)O-, -O(CO)O-, -N(R₆)-O-, -O-NR₅₁-, thio, sulfinyl, sulfonyl, -SO₂NR₇-, -NR₅₂SO₂- or -NR₈-;

 R_2 is a C_1 - C_8 alkyl, C_3 - C_6 alkenyl or C_3 - C_6 alkynyl group which is mono- or polysubstituted by halogen, hydroxyl, amino, formyl, nitro, cyano, mercapto, carbamoyl, C_1 - C_6 alkoxy, C_1 - C_6 alkoxycarbonyl, C_2 - C_6 alkenyl, C_2 - C_6 haloalkenyl, C_2 - C_6 alkynyl, C_2 - C_6 haloalkynyl, C_3 - C_6 cycloalkyl, by halogen-substituted C_3 - C_6 cycloalkyl, or by C_3 - C_6 alkenyloxy, C_3 - C_6 alkoxy, C_1 - C_6 alkoxy, C_3 - C_6 haloalkenyloxy, cyano- C_1 - C_6 alkoxy, C_1 - C_6 alkoxy, C_1 - C_6 alkoxy- C_1 - C_6 alkoxy- C_1 - C_6 alkoxy- C_1 - C_6 alkoxy, C_1 - C_6 alkoxycarbonyl, C_1 - C_6 alkylsulfonyl, C_1 - C_6 alkylsulfonyl, oxiranyl, which for its part may be substituted by C_1 - C_6 alkyl, or by benzylthio, benzylsulfonyl, benzylsulfonyl, C_1 - C_6 alkylamino, di(C_1 - C_6 alkyl)amino, C_1 - C_6 alkyl, or by benzylthio, benzylsulfonyl, benzylsulfonyl, phenoxy, phenylthio, phenylsulfinyl or phenylsulfonyl;

where the phenyl- or benzyl-containing groups for their part may be substituted by one or more C_1 - C_6 alkyl, C_1 - C_6 haloalkyl, C_1 - C_6 alkoxy, C_1 - C_6 haloalkoxy, halogen, cyano, hydroxyl or nitro groups, or

 R_2 is phenyl which may be mono- or polysubstituted by C_1 - C_6 alkyl, C_1 - C_6 haloalkyl, C_1 - C_6 alkoxy, C_1 - C_6 haloalkoxy, halogen, cyano, hydroxyl or nitro; or R_2 is C_3 - C_6 cycloalkyl, C_1 - C_6 alkoxyl- or C_1 - C_6 alkyl-substituted C_3 - C_6 cycloalkyl, 3-oxetanyl or C_1 - C_6 alkyl-substituted 3-oxetanyl;

or, if Q is Q2 or Q3, or is Q1 in which R14 and R22 are a C2-C3alkylene chain, R2 is additionally also a five- to ten-membered monocyclic or fused bicyclic ring system which may be aromatic, saturated or partially saturated and may contain 1 to 4 heteroatoms selected from the group consisting of nitrogen, oxygen and sulfur, where the ring system is attached directly or via a C₁-C₄alkylene, C₂-C₄alkenyl-C₁-C₄alkylene, C₂-C₄alkynyl-C₁-C₄alkylene, -N(R₁₂)-C₁-C₄alkylene, -SO-C₁-C₄alkylene or -SO₂-C₁-C₄alkylene group to the substituent X₁ and where each ring system may not contain more than two oxygen atoms and not more than two sulfur atoms and the ring system for its part may be mono-, di- or trisubstituted by C_1 - C_6 alkyl, C_1 - C_6 haloalkyl, C_2 - C_6 alkenyl, C_2 - C_6 haloalkynyl, C_2 - C_6 haloalkynyl, C₁-C₆alkoxy, hydroxyl, C₁-C₆haloalkoxy, C₃-C₆alkenyloxy, C₃-C₆alkynyloxy, mercapto, C₁-C₆alkylthio, C₁-C₆haloalkylthio, C₃-C₆alkenylthio, C₃-C₆haloalkenylthio, C₃-C₆alkynylthio. C_2 - C_5 alkoxyalkylthio, C_3 - C_5 acetylalkylthio, C_3 - C_6 alkoxycarbonylalkylthio, C_2 - C_4 cyanoalkylthio, C1-C6alkylsulfinyl, C1-C6haloalkylsulfinyl, C1-C6alkylsulfonyl, C1-C6haloalkylsulfonyl, aminosulfonyl, C1-C2alkylaminosulfonyl, di(C1-C2alkyl)aminosulfonyl, di(C1-C4alkyl)amino, halogen, cyano, nitro, phenyl and benzylthio, where phenyl and benzylthio for their part may be substituted on the phenyl ring by C₁-C₃alkyl, C₁-C₃haloalkyl, C₁-C₃alkoxy,

C₁-C₃haloalkoxy, halogen, cyano or nitro, and where the substituents on the nitrogen in the heterocyclic ring are different from halogen; or

R₂ is hydrogen or unsubstituted C₁-C₈alkyl if

- a) R₁ is substituted by the group R₅, or
- b) Q is the group Q₂, or
- c) Q is the group Q_3 in which X_1 is -O(CO)-, -(CO)O-, -N(R₆)-O-, -O-NR₅₁-, -SO₂NR₇-, -NR₅₂SO₂- or -NR₈-; or
- d) Q is the group Q_1 in which X_1 is -N(R₆)-O-, -O-NR₅₁-, -SO₂NR₇-, -NR₅₂SO₂- or -NR₈-, or
- e) Q is the group Q_1 in which R_{14} and R_{22} in Q_1 are a C_2 - C_3 alkylene chain and X_1 is -O(CO)-or -(CO)O-,

R₃ is C₁-C₃haloalkyl;

 R_4 is hydrogen, halogen, C_1 - C_3 alkyl, C_1 - C_3 haloalkyl, C_1 - C_3 alkoxy, C_1 - C_3 alkoxy- C_1 - C_3 alkoxy;

 R_5 is hydroxyl, C_1 - C_6 alkoxy, C_3 - C_6 cycloalkyloxy, C_1 - C_6 alkoxy- C_1 - C_6 alkoxy, C_1 - C_6 alkoxy or C_1 - C_2 alkylsulfonyloxy;

 R_6 , R_7 , R_8 , R_9 , R_{10} , R_{11} , R_{12} , R_{51} and R_{52} independently of one another are hydrogen, C_1 - C_6 alkyl, C_1 - C_6 alkyl, C_1 - C_6 alkoxy- C_1 - C_6 alkoxy, benzyl or phenyl, where phenyl and benzyl for their part may be mono- or polysubstituted by C_1 - C_6 alkyl, C_1 - C_6 haloalkyl, C_1 - C_6 alkoxy, C_1 - C_6 haloalkoxy, halogen, cyano, hydroxyl or nitro; where R_6 and R_9 are not simultaneously hydrogen and hydrogen, C_1 - C_6 alkoxycarbonyl or C_1 - C_6 alkylcarbonyl, respectively;

Q is Q₁

in which

 A_1 is $C(R_{14}R_{15})$, NR_{16} or oxygen;

 A_2 is $C(R_{17}R_{18})$, C(O), $-C=N-O-R_{19}$, oxygen, thio, sulfinyl, sulfonyl, $-NR_{20}$ or ethylene; with the provisos that A_1 is different from oxygen if A_2 is oxygen, C(O), thio, sulfinyl, $-C=N-O-R_{19}$, NR_{20} or $C(R_{17}R_{18})$, where R_{17} and R_{18} independently of one another are C_1-C_4 alkoxy, C_1-C_4 alkylsulfinyl, C_1-C_4 alkylsulfonyl; and A_1 is different from NR_{16} if A_2 is thio, sulfinyl or $C(R_{17}R_{18})$, where R_{17} and R_{18} independently of one another are C_1-C_4 alkoxy, C_1-C_4 alkylsulfinyl, C_1-C_4 alkylsulfonyl; R_{14} and R_{22} independently of one another are hydrogen, C_1-C_4 alkyl, C_1-C_4 alkylsulfonyl, C_3-C_4 alkenyl, C_3-C_4 alkynyl, C_1-C_4 alkylsulfonyl, C_1-C_4 al

C₁-C₄alkoxycarbonyl or C₁-C₄alkylcarbonyl;

 R_{15} and R_{21} independently of one another are hydrogen, C_1 - C_4 alkyl, C_1 - C_4 haloalkyl, C_3 - C_4 alkenyl or C_3 - C_4 alkynyl;

 R_{17} is hydrogen, C_1 - C_4 alkyl, C_1 - C_4 haloalkyl, C_1 - C_4 alkoxy, C_1 - C_4 alkylsulfonyl;

 R_{18} is hydrogen, C_1 - C_4 alkyl, C_1 - C_4 haloalkyl, C_3 - C_4 alkenyl, C_3 - C_4 alkynyl, C_1 - C_4 alkylsulfinyl, C_1 - C_4 alkylsulfonyl or C_1 - C_4 dialkoxyalkyl- C_1 - C_4 alkyl;

 R_{20} is C_1 - C_4 alkyl, C_3 - C_6 cycloalkyl, C_3 - C_4 alkenyl, C_3 - C_4 alkynyl, C_1 - C_4 alkylcarbonyl, C_1 - C_4 alkylcarbonyl, C_1 - C_4 alkylcarbonyl, carbonyloxy, di(C_1 - C_4 alkyl)aminocarbonyl or benzyl, where the phenyl group may be monoor polysubstituted by C_1 - C_6 alkyl, C_1 - C_6 haloalkyl, C_1 - C_6 alkoxy, C_1 - C_6 haloalkoxy, halogen, cyano, hydroxyl or nitro;

 R_{19} and R_{16} independently of one another are hydrogen, C_1 - C_4 alkyl, C_3 - C_6 cycloalkyl, C_3 - C_4 alkenyl, C_3 - C_4 alkenyl, benzyl or phenyl, where phenyl and benzyl for their part may be mono- or polysubstituted by C_1 - C_6 alkyl, C_1 - C_6 haloalkyl, C_1 - C_6 alkoxy, C_1 - C_6 haloalkoxy, halogen, cyano, hydroxyl or nitro;

or R₁₄ and R₂₂ together form a C₂-C₃alkylene chain;

or R_{14} and R_{15} together and/or R_{17} and R_{18} together and/or R_{21} and R_{22} together form a C_2 - C_4 alkylene chain which may be interrupted by oxygen and/or carbonyl and/or sulfur, with the proviso that the oxygen and sulfur atoms are separated by at least one methylene group; or R_{14} and R_{18} together form a C_2 - C_4 alkylene chain; or

R₂₂ and R₁₈ together form a C₂-C₄alkylene chain;

or R₁₈ together with R₂₂ or R₁₄ forms a direct bond;

or R₁₆ and R₁₈ together form a C₂-C₄alkylene chain;

R₁₃ is hydroxyl, O⁻M⁺, where M⁺ is an alkali metal cation or ammonium cation, halogen.

C₁-C₁₂alkylsulfonyloxy, amino, C₁-C₄alkylthio, C₁-C₁₂alkylsulfonyl, C₁-C₁₂alkylsulfonyl,

 C_1 - C_{12} haloalkylthio, C_1 - C_{12} haloalkylsulfinyl, C_1 - C_{12} haloalkylsulfonyl,

 $C_1-C_6 alkoxy-C_1-C_6 alkylsulfinyl, \ C_1-C_6 alkoxy-C_1-C_6 alkylsulfinyl, \ C_1-C_6 alkoxy-C_1-C_6 alkylsulfonyl, \ C_1-C_6 alkylsulfonyl, \$

 C_3 - C_{12} alkenylthio, C_3 - C_{12} alkenylsulfinyl, C_3 - C_{12} alkenylsulfonyl, C_3 - C_{12} alkynylthio,

 $C_3\text{-}C_{12} \\ \text{alkynylsulfinyl}, \ C_3\text{-}C_{12} \\ \text{alkynylsulfonyl}, \ C_1\text{-}C_4 \\ \text{alkoxycarbonyl-}C_1\text{-}C_4 \\ \text{alkylthio}, \ C_2\text{-}C_4 \\ \text{alkynylsulfonyl}, \ C_3\text{-}C_4 \\ \text{alkynylsulfonylsul$

 $C_1\text{-}C_4 alkoxy carbonyl-C_1\text{-}C_4 alkyl sulfinyl, \ C_1\text{-}C_4 alkoxy carbonyl-C_1\text{-}C_4 alkyl sulfonyl, \ C_2\text{-}C_4 alkoxy carbonyl-C_1\text{-}C_4 alkyl sulfonyl, \ C_2\text{-}C_4 alkoxy carbonyl-C_2\text{-}C_4 alkyl sulfonyl, \ C_2\text{-}C_4 alkoxy carbonyl-C_2\text{-}C_4 alkyl sulfonyl, \ C_2\text{-}C_4 alkoxy carbonyl-C_2\text{-}C_4 alkyl sulfonyl, \ C_2\text{-}C_4 alkyl sulfonyl,$

 $(C_1-C_4$ alkoxy)₂P(O)O, C_1-C_4 alkyl- $(C_1-C_4$ alkoxy)P(O)O, $H(C_1-C_4$ alkoxy)P(O)O,

 $R_{23}R_{24}NR_{25}R_{26}N$, NH, $R_{27}R_{28}NC(O)O$ -, $R_{29}R_{30}NC(O)NH$ -, C_1 - C_{18} alkylcarbonyloxy,

 C_2 - C_{18} alkenylcarbonyloxy, C_2 - C_{18} alkynylcarbonyloxy, C_3 - C_6 cycloalkylcarbonyloxy,

 C_1 - C_{12} alkoxycarbonyloxy, C_1 - C_{12} alkylthiocarbonyloxy, C_1 - C_{12} alkylthiocarbamoyl, where the alkyl, alkenyl and alkynyl groups may be substituted by halogen, C_1 - C_6 alkylthio, C_1 - C_6 alkylsulfonyl or cyano;

or R_{13} is phenoxy, phenylthio, phenylsulfinyl, phenylsulfonyl, phenylsulfonylamino, phenylsulfonyloxy or benzoyloxy, where the phenyl groups for their part may be substituted by one or more halogen, nitro, cyano, C_1 - C_4 alkyl, C_1 - C_4 haloalkyl, C_1 - C_4 alkoxy or C_1 - C_4 haloalkoxy groups;

or R₁₃ is a group Het₁-thio, Het₂-sulfinyl, Het₃-sulfonyl, Het₄-(CO)O or Het₅-N(R₃₃); in which

Het₁, Het₂, Het₃, Het₄ and Het₅ independently of one another are a five- to ten-membered monocyclic or fused bicyclic ring system which may be aromatic or partially saturated and may contain 1 to 4 heteroatoms selected from the group consisting of nitrogen, oxygen and sulfur, and where each ring system may not contain more than 2 oxygen atoms and not more than 2 sulfur atoms, and where the ring system itself can be substituted by C₁-C₆alkyl, C₁-C₆haloalkyl, C₁-C₆alkoxy, C₁-C₆haloalkoxy, C₁-C₆alkylthio, C₁-C₆alkylsulfinyl, C₁-C₆alkylsulfonyl, di(C₁-C₄alkyl)aminosulfonyl, di(C₁-C₄alkyl)amino, halogen, cyano, nitro or phenyl, and where the substituents on the nitrogen in the heterocyclic ring are different from halogen;

R₂₃, R₂₄, R₂₅, R₂₆, R₂₇, R₂₈, R₂₉, R₃₀ and R₃₃ independently of one another are hydrogen or C₁-C₆alkyl;

or R_{23} and R_{24} together or R_{25} and R_{26} together or R_{27} and R_{28} together or R_{29} and R_{30} together are pyrolidino, piperidino, morpholino, thiomorpholino, which may be mono- or polysubstituted by methyl groups;

or Q is Q2

in which

R₃₄ is hydrogen, C₁-C₄alkyl, C₁-C₄haloalkyl, C₃-C₆cycloalkyl, C₂-C₄alkenyl, C₂-C₄alkynyl or benzyl, where the phenyl group may be mono- or polysubstituted by C1-C6alkyl, C1-C6haloalkyl, C₁-C₆alkoxy, C₁-C₆haloalkoxy, halogen, cyano, hydroxyl or nitro; R₃₅ is hydrogen, C₁-C₄alkyl, C₁-C₄haloalkyl, C₃-C₆cycloalkyl, C₃-C₄alkenyl, C₃-C₄alkynyl or benzyl, where the phenyl group may be mono- or polysubstituted by C1-C6alkyl, C1-C6haloalkyl, C₁-C₆alkoxy, C₁-C₆haloalkoxy, halogen, cyano, hydroxyl or nitro; R₃₆ is hydroxyl, O'M⁺, where M⁺ is an alkali metal cation or ammonium cation, halogen,

C₁-C₁₂alkylsulfonyloxy, amino, C₁-C₄alkylthio, C₁-C₁₂alkylsulfinyl, C₁-C₁₂alkylsulfonyl,

C₁-C₁₂haloalkylthio, C₁-C₁₂haloalkylsulfinyl, C₁-C₁₂haloalkylsulfonyl,

 C_1 - C_6 alkoxy- C_1 - C_6 alkylthio, C_1 - C_6 alkoxy- C_1 - C_6 alkylsulfonyl, C_1 - C_6 alkoxy- C_1 - C_6 alkylsulfonyl, C₃-C₁₂alkenylthio, C₃-C₁₂alkenylsulfinyl, C₃-C₁₂alkenylsulfonyl, C₃-C₁₂alkynylthio,

C₃-C₁₂alkynylsulfinyl, C₃-C₁₂alkynylsulfonyl, C₁-C₄alkoxycarbonyl-C₁-C₄alkylthio,

C₁-C₄alkoxycarbonyl-C₁-C₄alkylsulfinyl, C₁-C₄alkoxycarbonyl-C₁-C₄alkylsulfonyl,

 $(C_1-C_4alkoxy)_2P(O)O$, $C_1-C_4alkyl-(C_1-C_4alkoxy)P(O)O$, $H(C_1-C_4alkoxy)P(O)O$, $R_{37}R_{38}N$, $R_{39}R_{40}NNH$, $R_{41}R_{42}NC(O)O$ -, $R_{43}R_{44}NC(O)NH$ -, $C_1-C_{18}alkylcarbonyloxy$, $C_2-C_{18}alkynylcarbonyloxy$, $C_3-C_6cycloalkylcarbonyloxy$, $C_1-C_{12}alkylthiocarbonyloxy$, $C_1-C_{12}alkylthiocarbonyloxy$, $C_1-C_{12}alkylthiocarbonyloxy$, where the alkyl, alkenyl and alkynyl groups may be substituted by halogen, $C_1-C_6alkoxy$, $C_1-C_6alkylthio$,

 R_{36} is phenoxy, phenylthio, phenylsulfinyl, phenylsulfonyl, phenylsulfonylamino, phenylsulfonyloxy or benzoyloxy, where the phenyl groups for their part may be mono- or polysubstituted by halogen, nitro, cyano, C_1 - C_4 alkyl, C_1 - C_4 haloalkyl, C_1 - C_4 alkoxy, C_1 - C_4 haloalkoxy,

C₁-C₆alkylsulfinyl, C₁-C₆alkylsulfonyl or cyano; or

or R₃₆ is a group Het₇-thio, Het₈-sulfinyl, Het₉-sulfonyl, Het₁₀-(CO)O or Het₁₁-N(R₄₇); in which Het₇, Het₈, Het₉, Het₁₀ and Het₁₁ independently of one another are a five- to ten-membered monocyclic or fused bicyclic ring system which may be aromatic or partially saturated and may contain 1 to 4 heteroatoms selected from the group consisting of nitrogen, oxygen and sulfur, and where each ring system may not contain more than 2 oxygen atoms and not more than 2 sulfur atoms, and where the ring system for its part may be substituted by C₁-C₆alkyl, C₁-C₆haloalkyl, C₁-C₆haloalkoxy, C₁-C₆haloalkoxy, C₁-C₆alkylthio, C₁-C₆alkylsulfinyl, C₁-C₆alkylsulfonyl, di(C₁-C₄alkyl)aminosulfonyl, di(C₁-C₄alkyl)amino, halogen, cyano, nitro or phenyl, and where the substituents on the nitrogen in the heterocyclic ring are different from halogen;

 R_{37} , R_{38} , R_{39} , R_{40} , R_{41} , R_{42} , R_{43} , R_{44} and R_{47} independently of one another are hydrogen or C_1 - C_6 alkyl; or

 R_{37} and R_{38} together or R_{39} and R_{40} together or R_{41} and R_{42} together or R_{43} and R_{44} together are pyrolidino, piperidino, morpholino, thiomorpholino, which may be mono- or polysubstituted by methyl groups; or Q is Q_3

$$R_{50}$$
-S(O)_n R_{49} (Q_3) ;

in which

 R_{49} is C_1 - C_4 alkyl, C_1 - C_4 haloalkyl, C_3 - C_6 cycloalkyl or halogen-substituted C_3 - C_6 cycloalkyl;

 R_{50} is C_1 - C_3 alkylene which may be substituted by halogen, hydroxyl, C_1 - C_6 alkoxy, C_2 - C_6 alkenyl, C_2 - C_6 alkynyl, C_3 - C_6 cycloalkyl, C_1 - C_6 alkoxy- C_1 - C_6 alkoxy, C_1 - C_6 alkoxy- C_1 - C_6 alkoxy- C_1 - C_6 alkoxy, (3-oxetanyl)oxy, or by C_1 - C_6 alkyl-substituted (3-oxetanyl)oxy, or by benzylthio, benzylsulfinyl, benzylsulfonyl, phenyl, phenoxy, phenylthio, phenylsulfinyl or phenylsulfonyl, where the phenyl- and benzyl-containing groups for their part may be substituted by one or more C_1 - C_6 alkyl, C_1 - C_6 haloalkyl, C_1 - C_6 alkoxy, C_1 - C_6 haloalkoxy, halogen, cyano, hydroxyl or nitro groups;

or R₅₀ is phenyl, where the phenyl-containing group for its part may be substituted by one or more C₁-C₆alkyl, C₁-C₆haloalkyl, C₁-C₆alkoxy, C₁-C₆haloalkoxy, halogen, cyano, hydroxyl or nitro,

or R_{50} is C_3 - C_6 cycloalkyl, C_1 - C_6 alkoxy- or C_1 - C_6 alkyl-substituted C_3 - C_6 cycloalkyl, 3-oxetanyl or C_1 - C_6 alkyl-substituted 3-oxetanyl; and

n is 0, 1 or 2; and agronomically acceptable salts/N-oxides/isomers/enantiomers of this compound.

2. A process for preparing compounds of the formula I as claimed in claim 1, which comprises, to prepare compounds of the formula I, in which R_1 , R_2 , R_3 , R_4 and X_1 are as defined under formula I and Q is a group Q_1 , either a) reacting a compound of the formula Ia

$$R_4$$
 R_3
 N
 R_1
 X_1
 R_2
 $(Ia),$

in which R_1 , R_2 , R_3 , R_4 and X_1 are as defined under formula I and Y_1 is a leaving group, in an inert organic solvent in the presence of a base with a compound of the formula II

in which R_{22} , R_{21} , A_2 and A_1 are as defined under formula I, to give the compounds of the formulae IIIa and IIIb

$$R_4$$
 R_3
 R_1
 R_2
 R_3
 R_1
 R_2
 R_3
 R_4
 R_3
 R_4
 R_3
 R_4
 R_5
 R_5

and then isomerizing these in the presence of a base and a catalytic amount of dimethylaminopyridine (DMAP) or a source of cyanide, for example acetone cyanohydrin; or b) reacting a compound of the formula lb

$$R_4$$
 OH R_3 N R_1 X_1 R_2 (lb),

in which R_1 , R_2 , R_3 , R_4 and X_1 are as defined under formula I, with a compound of the formula II

in which R_{22} , R_{21} , A_1 and A_2 are as defined under formula I, in an inert organic solvent in the presence of a base and a coupling agent to give the compound of the formula IIIa or IIIb,

and then isomerizing these as described under route a); or, to prepare the compounds of the formula I, in which Q is a group Q_2 , either a) reacting a compound of the formula Ia

$$R_3$$
 N R_1 X_1 R_2 (Ia),

in which R_1 , R_2 , R_3 , R_4 and X_1 are as defined under formula I and Y_1 is a leaving group, with a compound of the formula IIa

$$R_{34}$$
 N
 N
 O
 R_{35}
(IIa),

in which R_{34} and R_{35} are as defined under formula I, in an inert organic solvent in the presence of a base to give the compound of the formula IIIc

$$R_{34}$$
 R_{35}
 R_{35}
 R_{35}
 R_{35}
 R_{35}
 R_{35}

in which R_1 , R_2 , R_3 , R_4 , R_{34} , R_{35} and X_1 are as defined under formula I, and then isomerizing this compound in the presence of a base and a catalytic amount of a source of cyanide; or

b) reacting a compound of the formula lb

$$R_4$$
 OH OH R_3 N R_1 X_1 R_2 (Ib),

in which $R_1,\,R_2,\,R_3,\,R_4$ and X_1 are as defined under formula I, with a compound of the formula IIa

in which R_{34} and R_{35} are as defined above, in an inert organic solvent in the presence of a base and a coupling agent to give the compound of the formula IIIc

$$R_{4}$$
 R_{3}
 R_{3}

and then isomerizing this compound as described under route a); or, to prepare compounds of the formula I, in which ${\sf Q}$ is a group

in which n is 0 and R_{50} and R_{49} are as defined above, either a) converting a compound of the formula IV

$$R_4$$
 R_3
 R_1
 X_1
 R_2
(IV),

in which X_1 , R_1 , R_2 , R_3 , R_4 and R_{49} are as defined above, in the presence of a base, carbon disulfide and an alkylating agent of the formula V

$$R_{50}-Y_2$$
 (V),

in which R_{50} is as defined under formula I, and Y_2 is a leaving group, into the compound of the formula VI

in which R_1 , R_2 , R_3 , R_4 , R_{50} , X_1 and R_{49} are as defined under formula I, and then cyclizing this compound with hydroxylamine hydrochloride in the presence of a base to give the isomeric compounds of the formulae Ic and Id

$$\begin{array}{c} \text{(O)} \text{n} & \text{R}_{50} \\ \text{N} \\ \text{R}_{3} & \text{N} \\ \text{R}_{1} \\ \text{X}_{1} \\ \text{R}_{2} \\ \text{(Ic)} \end{array} \quad \text{and} \quad \begin{array}{c} \text{(O)} \text{n} & \text{R}_{50} \\ \text{O} \\ \text{N} \\ \text{R}_{49} \\ \text{R}_{3} \\ \text{N} \\ \text{R}_{1} \\ \text{R}_{2} \\ \text{(Id)} \end{array}$$

and then oxidizing these compounds with an oxidizing agent, for example with peracids, to give the corresponding sulfoxides (n = 1) and sulfones (n = 2) of the formulae le and If, respectively.

- 3. A herbicidal and plant-growth-inhibiting composition, which contains a herbicidally effective amount of a compound of the formula I on an inert carrier.
- 4. A method for controlling undesirable plant growth, wherein a herbicidally effective amount of an active compound of the formula I or a composition which contains this active compound is applied to the plants or their habitat.
- 5. A method for inhibiting plant growth, wherein a herbicidally effective amount of an active compound of the formula I or a composition which contains this active compound is applied to the plants or their habitat.
- 6. A compound of the formula XX

in which

Q is hydroxyl, halogen, cyano or C₁-C₆alkoxy, or is a group of the formula

R₁, R₃, R₄, R₄₉, R₅₀, X₁ and p are as defined under formula I and R₂ is a C₁-C₈alkyl, C₃-C₆alkenyl or C₃-C₆alkynyl group which is mono- or polysubstituted by halogen, hydroxyl, amino, formyl, nitro, cyano, mercapto, carbamoyl, C1-C6alkoxy, C1-C6alkoxycarbonyl, C2-C6alkenyl, C2-C6haloalkenyl, C2-C6alkynyl, C2-C6haloalkynyl, C3-C6cycloalkyl, by halogensubstituted C₃-C₆cycloalkyl, or by C₃-C₆alkenyloxy, C₃-C₆alkynyloxy, C₁-C₆haloalkoxy, C₃-C₆haloalkenyloxy, cyano-C₁-C₆alkoxy, C₁-C₆alkoxy-C₁-C₆alkoxy, C₁-C₆alkoxy-C₁-C₆alkoxy-C₁-C₆alkoxy, C₁-C₆alkylthio-C₁-C₆alkoxy, C₁-C₆alkylsulfinyl-C₁-C₆alkoxy, C₁-C₆alkylsulfonyl-C₁-C₆alkoxy, C₁-C₆alkoxycarbonyl-C₁-C₆alkoxy, C₁-C₆alkoxycarbonyl, C₁-C₆alkylcarbonyl, C₁-C₆alkylthio, C₁-C₆alkylsulfinyl, C₁-C₆alkylsulfonyl, C₁-C₆haloalkylthio, C1-C6-haloalkylsulfinyl, C1-C6-haloalkylsulfonyl, oxiranyl, which for its part may be substituted by C₁-C₆alkyl, or by (3-oxetanyl)oxy, which for its part may be substituted by C₁-C₆alkyl, or by benzylthio, benzylsulfinyl, benzylsulfonyl, C1-C6alkylamino, di(C1-C6alkyl)amino, R9S(O)2O, R₁₀N(R₁₁)SO₂-, thiocyanato, phenyl, phenoxy, phenylthio, phenylsulfinyl or phenylsulfonyl; where the phenyl- or benzyl-containing groups for their part may be substituted by one or more C_1 - C_6 alkyl, C_1 - C_6 haloalkyl, C_1 - C_6 alkoxy, C_1 - C_6 haloalkoxy, halogen, cyano, hydroxyl or nitro groups, or

 R_2 is phenyl which may be mono- or polysubstituted by C_1 - C_6 alkyl, C_1 - C_6 haloalkyl, C_1 - C_6 alkoxy, C_1 - C_6 haloalkoxy, halogen, cyano, hydroxyl or nitro; or R_2 is C_3 - C_6 cycloalkyl, C_1 - C_6 alkoxy- or C_1 - C_6 alkyl-substituted C_3 - C_6 cycloalkyl, 3-oxetanyl or C_1 - C_6 alkyl-substituted 3-oxetanyl;

or

if X_1 is -N(R₆)-O-, -O-NR₅₁, SO₂NR₇- or -NR₅₂SO₂- and R₆, R₇, R₅₁ and R₅₂ are as defined under formula I,

 R_2 may additionally be hydrogen, unsubstituted C_1 - C_6 alkyl, or a five- to ten-membered monocyclic or fused bicyclic ring system which may be aromatic, saturated or partially saturated and may contain 1 to 4 heteroatoms selected from the group consisting of nitrogen, oxygen and sulfur, where the ring system is attached directly or via a

 $C_1\text{-}C_4\text{alkylene, } C_2\text{-}C_4\text{alkenyl-}C_1\text{-}C_4\text{alkylene, } C_2\text{-}C_4\text{alkylene, } C_2\text{-}C_4\text{-}C_4\text{alkylene, } C_2\text{-}C_6\text{-}C_4\text{alkylene, } C_2\text{-}C_6\text{-}C_4\text$

INTERNATIONAL SEARCH REPORT

Inter "nat Application No PC I/ EP 01/06430

A. CLASSI IPC 7	FICATION OF SUBJECT MATTER C07D401/06 C07D405/06 C07D409, C07D213/50 A01N43/40	/06 C07D413/06	C07D417/06
	o International Patent Classification (IPC) or to both national classific SEARCHED	alion and IPC	· ·
	ocumentation searched (classification system followed by classification CO7D AO1N	ion symbols)	
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	ENTS CONSIDERED TO BE RELEVANT		
Category °	Citation of document, with indication, where appropriate, of the re	levant passages	Relevant to claim No.
X,P	WO 00 39094 A (NOVARTIS AG.,	-	1-6
	SWITZ.;NOVARTIS-ERFINDUNGEN VERWALTUNGSGESELLSCHAFT M.)		
	6 July 2000 (2000-07-06)		·
	cited in the application claim 1		
χ	 ₩O 00 15615 A (NOVARTIS AG.,		1-6
^	SWITZ.; NOVARTIS-ERFINDUNGEN		1-0
	VERWALTUNGSGESELLSCHAFT M.) 23 March 2000 (2000-03-23)		
	cited in the application		
	claim 1		
A	US 5 260 262 A (LEE, LEN FANG ET 9 November 1993 (1993-11-09)	AL)	1-6
	claim 1		
			·
	ner documents are listed in the continuation of box C.	X Patent family members are	e listed in annex.
	tegories of cited documents:	*T* later document published after to or priority date and not in confi	he international filing date lict with the application but
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which i citation	is cited to establish the publication date of another or other special reason (as specified)	"Y" document of particular relevance cannot be considered to involve	e; the claimed invention e an inventive step when the
other n	-	document is combined with on- ments, such combination being in the art.	e or more other such docu-
tater th	nt published prior to the international filing date but an the priority date claimed	*&* document member of the same	patent family
Date of the	actual completion of the international search	Date of mailing of the internation	onal search report
23	3 July 2001	31/07/2001	
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INTERNATIONAL SEARCH REPORT

Inte mal Application No PCTEP 01/06430

	atent document d in search report	ı.	Publication date	1	Patent family member(s)	Publication date
WO	0039094	Α	06-07-2000	AU	2101500 A	31-07-2000
WO	0015615	A	23-03-2000	AU BR EP	5862999 A 9913745 A 1114030 A	03-04-2000 05-06-2001 11-07-2001
US	5260262	A	09-11-1993	AT AU CA DE DE EP ES JP US WO US	184596 T 3221293 A 2122262 A 69230003 D 69230003 T 0621863 A 2137249 T 7502993 T 5298479 A 9311112 A 5380699 A	15-10-1999 28-06-1993 10-06-1993 21-10-1999 30-03-2000 02-11-1994 16-12-1999 30-03-1995 29-03-1994 10-06-1993 10-01-1995